

Velocity control in Parkinson's disease: a quantitative analysis of isochrony in scribbling movements

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Abstract An experiment was conducted to contrast the motor performance of three groups ($N = 20$) of participants: (1) patients with confirmed Parkinson Disease (PD) diagnose; (2) age-matched controls; (3) young adults. The task consisted of scribbling freely for 10 s within circular frames of different sizes. Comparison among groups focused on the relation between the figural elements of the trace (overall size and trace length) and the velocity of the drawing movements. Results were analysed within the framework of previous work on normal individuals showing that instantaneous velocity of drawing movements depends jointly on trace curvature (Two-thirds Power Law) and trace extent (Isochrony principle). The motor behaviour of PD patients exhibited all classical symptoms of the disease (reduced average velocity, reduced fluency, micrographia). At a coarse level of analysis both isochrony and the dependence of velocity on curvature, which are

supposed to reflect cortical mechanisms, were spared in PD patients. Instead, significant differences with respects to the control groups emerged from an in-depth analysis of the velocity control suggesting that patients did not scale average velocity as effectively as controls. We factored out velocity control by distinguishing the influence of the broad context in which movement is planned—i.e. the size of the limiting frames—from the influence of the local context—i.e. the linear extent of the unit of motor action being executed. The balance between the two factors was found to be distinctively different in PD patients and controls. This difference is discussed in the light of current theorizing on the role of cortical and sub-cortical mechanisms in the aetiology of PD. We argue that the results are congruent with the notion that cortical mechanisms are responsible for generating a parametric template of the desired movement and the BG specify the actual spatio-temporal parameters through a multiplicative gain factor acting on both size and velocity.

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Introduction

Parkinson's disease (PD) is one of the most common and disabling neurodegenerative conditions in the general population that affects mainly the motor system. PD results from degenerative processes involving selectively, although not exclusively, the *Substantia Nigra pars compacta* (SNc). This leads to a progressive loss of the dopaminergic innervations of the striatum in the basal ganglia (BG) and to the cardinal symptoms of PD, which include tremor, rigidity, akinesia, postural abnormalities

and gait impairment. In particular, akinesia encompasses a variety of subtle but highly interfering motor disturbances that can be separated into distinct components including delay in movement initiation (true akinesia), slowing of ongoing movements (bradykinesia), reduction of movement amplitude (hypokinesia), fatigue during repetitive movements, paucity of spontaneous movements, and a reduced ability to produce sequences of alternating or complex movements. These distinct yet related aspects of akinesia usually occur in various combinations in individual patients and concur to produce many typical symptoms of PD such as amimia, hypophonia, dysarthria, micrographia, slowness of gait with reduced arm swing, and global immobility.

Motor deficits in PD

PD-related motor deficits have been analysed most extensively in single, goal-directed movements, which exhibit a well-established pattern of abnormalities. First, simple reaction times (RT) are longer in patients than in age-matched controls (Flowers 1975; Bloxham et al. 1984; Sheridan et al. 1987; Flash et al. 1992; Jahanshahi et al. 1992). Second, most patients show bradykinesia in both upper (Hallett and Khoshbin 1980; Ghilardi et al. 2000) and lower limbs (Morris et al. 1994; Morris et al. 2001). It is generally agreed that bradykinesia is caused by the inability to generate the required level of force at the appropriate rate (Godaux et al. 1992; Weiss et al. 1997; Berardelli et al. 2001). It is still debated, however, whether the problem is due to under-scaling of the central motor commands (Berardelli et al. 2001), or inability of the neuromuscular system to adapt quickly to the required force level (Weiss et al. 1997). Third, visual targets are systematically undershot (hypokinesia), with a tendency for the deficit to increase with target distance (Flowers 1975; Vinter and Gras 1998; Desmurget et al. 2003; Desmurget et al. 2004a). Consequently, PD patients also show an abnormally large range effect (Brown et al. 1948). Hypokinesia is most obvious in pointing hand movements and in handwriting (micrographia) (McLennan et al. 1972), but emerges also in gait control. Prompted to increase their walking speed, PD patients increase stepping rhythm much more than stride length (Morris et al. 1994), a poorly effective strategy leading to festination and repeated falls.

The tendency to undershoot visual targets is often associated with, but not a direct consequence of bradykinesia. The ability to estimate visually hand and target position in space appears to be preserved in PD patients (Desmurget et al. 2000, 2004a). The factors most likely responsible for the poor reaching performance in these patients include (1) incorrect timing of the motor

commands during ballistic movement (Hallett et al. 1977; Pfann et al. 2001); (2) altered kinaesthetic perception of the initial hand position (Vindras and Viviani 1998); (3) erroneous estimation of movement extent, which may result from a combination of increased reafferent signals and reduced effectiveness of the corollary discharge (Moore 1987, 1989); (4) ineffective use of proprioceptive information, both per se (Schneider et al. 1987; Demirci et al. 1997; Zia et al. 2000) and in conjunction with visual information (Adamovich et al. 2001; Desmurget et al. 2003).

Supplying appropriate visual or acoustic stimuli reduces all motor deficits mentioned above (with the possible exception of the RT increase)—a phenomenon known as paradoxical movement (Georgiou et al. 1993). For instance, PD patients are as fast as age-matched controls when they have to catch a moving ball, but much slower in reaching a stable target (Majsak et al. 1998). Similarly, PD patients exhibit almost normal stride length when the steps are guided by transversal white stripes on the floor, whereas in the absence of visual cues, initiation of gait is delayed leading to short, ineffective steps and freezing episodes (Morris et al. 1994). Several mechanisms have been suggested to explain the beneficial role of these stimuli. They include promoting access to motor programmes (Bloxham et al. 1984), activating cerebellar circuits (Glickstein and Stein 1991), and boosting the activation of the medial cortical area by BG (Berardelli et al. 2001).

What do PD symptoms tell about movement control?

The neurophysiological bases of PD and the associated motor symptoms strongly suggest that BG play an important role in the normal control of amplitude and velocity of voluntary movements. Neurophysiological (Turner and Anderson 1997) and brain imaging studies (Turner et al. 1998; Desmurget et al. 2004b) provide additional evidence along the same line. However, the specific contribution of the BG in normal individuals is still open to debate. At present, at least two views are being contrasted, which postulate a “scaling” (Wichmann and DeLong 1996), and a “focussing” (Mink 1996) role for BG, respectively. Both models agree that the cause of PD is the loss of neurones in the SNc, and the consequent reduction of dopaminergic drive to the striatum. The decrease has opposite consequences on the direct and indirect pathways from the striatum to the output nuclei of the BG, namely the *Globus Pallidus internal segment* (Gpi) and the *Substantia Nigra pars reticulata* (SNr). The activity in the direct inhibitory pathway is reduced, whereas the activity in the indirect excitatory pathway through the *Globus Pallidus external segment* (GPe) and the subthalamic nucleus is increased.

The two models diverge instead as far as the function of the neurons targeted by direct and indirect pathways to GPI and SNr neurons is concerned.

According to the “scaling” model, direct and indirect pathways are activated sequentially. First, by inhibiting the GPI/SNr the direct pathway would decrease their inhibitory output and, therefore, facilitate movement. The indirect pathway would be activated after a delay to terminate the ongoing movement. Specifically, the activation would relieve inhibition of GPI/SNr and therefore increase the inhibition of the thalamic and brainstem neurones onto which GPI/SNr project. According to this model, the BG would play both a gating role by sharpening onset and offset phases of the movement, and a scaling role by modulating force.

The second (“focussing”) model holds instead that both direct and indirect pathways projecting on GPI/SNr neurones would simultaneously disinhibit the target motor centres and, at the same time, inhibit competing motor mechanism that would otherwise interfere with the desired movement. Thus, PD would prevent both the removal of inhibition from the desired motor pattern generator and the inhibition of the competing patterns.

In addition, it has been suggested that BG contribute to movement execution by modulating the spinal reflex circuits via their descending output to the brainstem (Johnson et al. 1991; Baldissera et al. 1994; Munro-Davies et al. 1999; Braak et al. 2000; Delwaide et al. 2000; Meunier et al. 2000).

It is still unclear whether PD-related motor deficits reflect a difficulty in the planning phase, associated with cortical dysfunction (Schwab et al. 1954; Marsden 1982; Berardelli et al. 2001; Desmurget et al. 2004b), or arise at the level of movement execution (Weiss et al. 1997), possibly because of associated brainstem disorders (Delwaide et al. 2000; Meunier et al. 2000; Pahapill and Lozano 2000; Simonetta-Moreau et al. 2002; Braak et al. 2003). Most arguments in support of one view or the other have been countered by different arguments. In favour of the planning hypothesis, the fact has been cited that PD patients engaged in a tracking task cannot take advantage of the predictive nature of the target as normal individuals do (Flowers 1978a, b). However, subsequent studies (Bloxham et al. 1984; Day et al. 1984; Schnider et al. 1995) have shown that the deficit is not always present. Moreover, PD patients are able to increase movement velocity with target distance (Berardelli et al. 1986a, 1986b; see, however, Draper and Johns 1964; Flowers 1976; Warabi et al. 1986; Pfann et al. 2001), and comply with time constraints (Teasdale et al. 1990; Pfann et al. 2001). Finally, the directional modulation of kinematic parameters associated with the anisotropy of arm inertia is similar in healthy subjects and PD patients (Gordon et al.

1994b; Ghilardi et al. 2000), suggesting a preserved ability to control independently movement direction and amplitude.

The fact that RTs are longer in PD patients than in controls might signal a delayed activation of the required motor programs. However, this inference is also controversial because the difference between choice and simple RTs is similar in both populations (Evarts et al. 1981). In fact, a slow build-up of motor excitability might well be responsible for increasing the response delay (Pascual-Leone et al. 1994; Berardelli et al. 2001).

Performing simultaneous movements is close to impossible for PD patients (Benecke et al. 1986; Benecke et al. 1987; Castiello and Bennett 1997; see, however, Brown and Jahanshahi 1998), which certainly points to a limited planning capacity. Yet this limitation may also arise because attempts to compensate BG deficiencies by vicarious cortical processes tax too much the computational resources of the system (Berardelli et al. 2001).

Finally, the fact that PD patients tend to make frequent, long pauses in tasks such as drawing geometrical figures (Berardelli et al. 1986a, 1986b), pointing to visual targets (Weiss et al. 1997), and making alternating hand movements has often been ascribed to a planning deficit (Harrington and Haaland 1991), or to the inability to switch from one motor program to another (Benecke et al. 1987). However, even in this case the argument is not watertight because a problem with the implementation stage (e.g. the inability to switch rapidly between different force levels) may lead to very similar deficits (Berardelli et al. 1986a, 1986b; Weiss et al. 1997).

Very recently, Mazzoni et al. (2007) have suggested yet another interpretation of the generalized movement slowing induced by PD. By focussing on reaching movements, these authors have argued that bradykinesia is not the consequence of an impaired speed-accuracy trade-off because, with adequate training, patients can perform as accurately and as fast as control. Rather, PD would alter the balance between the perceived reward of arriving to the target quickly and the amount of energy or effort required to achieve the fast movement. In other words, fast and accurate movements are within the available choices for patients, but the disease makes their selection less likely because they are perceived as more costly energetically by patients than by controls. In line with this interpretation, the authors also suggest that striatal dopamine exerts an “energizing” action on a “motor motivational” system that governs automatic and spontaneous behaviour.

Insofar as it is generally agreed that the relation between movement extent and duration pertains more directly to the planning phase of the movement, an in-depth comparative analysis of this relation may afford a new window on the nature of the PD-related motor deficits and contribute to the

planning versus execution debate evoked above. The next section provides the background and the rationale for our specific approach to such a comparative analysis.

Isochrony

Previous works demonstrated that instantaneous movement velocity depends jointly on the curvature of the trajectory—via the so-called Two-thirds Power Law (Viviani and Terzuolo 1982; Lacquaniti et al. 1983)—and on the metric properties of the trajectory—via the so-called Isochrony Principle (Viviani and Terzuolo 1980; Viviani and McCollum 1983; Viviani and Zanone 1988; Viviani and Schneider 1991). Whereas the Two-thirds Power Law describes the local, instantaneous variations of velocity, the principle of isochrony refers to a global regularity present in many end-point movements (Lacquaniti et al. 1984), most clearly in drawing and writing gestures. The principle states that the average velocity with which the gesture is executed increases spontaneously as a function of its size, so that execution time is less dependent on size than it would be otherwise. Moreover, it has been shown (Ostry et al. 1987) that the shape of the velocity curve remains almost invariant across changes in movement duration. This general statement, however, needs to be qualified according to the type of movement involved.

For simple point-to-point (not necessarily straight) movements, a power function $V_m = K \times L^\alpha$ describes accurately the relation between the average velocity from onset to end (V_m) and the linear extent (L) of the trajectory (Freeman 1914; Michel 1971; Viviani and Terzuolo 1983; Viviani and McCollum 1983; Lacquaniti et al. 1984). Experimental estimates of the exponent α range between 0.80 and 1.00 indicating a very high degree of velocity compensation. For more complex movements that can be construed as a sequence of simpler units the principle applies both globally and piece-wise. For instance, in the case of cursive handwriting the average tracing velocity of each letter depends on the relative size of the letter, as well as on the size of the entire word (Lacquaniti et al. 1984). When the movement is periodic, i.e. when the same closed pattern is traced repeatedly, the average velocity covaries with the linear extent of the trajectory over one movement cycle (Viviani and McCollum 1983). Whenever the pattern can be decomposed into identifiable figural units, the isochrony equation for cyclic movements takes a more complex form. For instance, if one draws continuously a knotted double ellipse the average velocity within each ellipse is a function of both its own perimeter and of the sum of the two perimeters (Viviani and Cenzato 1985).

The findings summarized above suggest that the mechanism underlying isochrony has access to an internal representation of the motor plan that includes at least an

estimate of the linear extent of the to-be-executed trajectory. Moreover, whenever the gesture involves a sequence of constituent units of motor action, it must be assumed that the linear extent of each unit is equally available. Clearly, this does not imply that the linear extent is an explicit parameter of the motor plan (Viviani and Flash 1995). In fact it says little about the nature of the motor representation other than the control of velocity draws from some sort of knowledge about the length of the trajectory *before* it is executed. This logical inference seems to be challenged by the observation that a form of isochrony is present even when we scribble (Viviani 1986). By definition, no advance plan is available for these extemporaneous movements that can continue indefinitely. Insofar as the notion of total linear extent of the trajectory becomes meaningless for scribbles, any regularity emerging at the global level in the control of velocity either reflects an underlying regularity at the local level, i.e. at the level of the units of motor action that are planned and executed sequentially while we scribble, or implies the existence of a more global control parameter. It is not known which one of these two hypotheses accounts best for the observed behaviour. In either case, isochrony in scribbles would still emerge from the planning phase. However, insofar as the BG have been credited with a role in taking into account the context in which the movement is executed (Houk and Wise 1995), and in controlling its scale (Turner et al. 1998), evidence that scribbling velocity depends in a lawful manner on some global size parameter of the movement would invite the inference that processes beyond the motor cortex are involved in this form of isochrony. Because BG functions are certainly impaired by PD, contrasting how velocity in PD patients and normal individuals is controlled, both locally and globally, when a complete plan of the movement is not available in advance should be interesting on two counts. It may contribute to the planning versus execution debate on the origin of PD-related motor deficits. Moreover, the pattern of differences in the two populations should be conducive to clarify the more general issue of how cortical and BG mechanisms cooperate to control simultaneously direction, extent and velocity of hand movements. The aim of our study is to provide such a comparative study based on a relatively large population of participants.

Methods

Experimental population

Three groups of individuals ($N = 20$) participated to the experiment. The first group (PK) included PD patients (13 males, 7 females) selected among those reporting at the

Department of Neurology of the Geneva University Hospitals. All but one patient were right-handed. All patients had typical, asymmetric, levodopa-responsive PD, according to the United Kingdom PD Society Brain Bank criteria. At the time of testing, patients were being treated with individualized regimens of standard and slow-release levodopa/carbidopa or levodopa/benserazide, COMT inhibitors, dopamine agonists and amantadine. Disease severity ranged from Hoehn & Yahr stage 1.5–3 (Hoehn and Yahr 1967). The second group (Age-Matched Controls; AMC) included individuals (14 males, 6 females, all right-handed) with no known neurological disorder. Table 1 reports detailed personal data for these two groups. Statistical analysis (*t* test) showed no significant age difference between the groups (PK: mean = 68.3, SD = 8.0; AMC: mean = 66.9, SD = 6.8) that were tested at the Department of Psychology of the University of Geneva. The third group (Young Controls: YC) included students (7 males, 13 females, all right-handed) of the Faculty of Psychology of UHSR University whose age ranged between 21 and 28 years. They were tested at Laboratory of Action, Perception and Cognition of the UHSR Faculty of Psychology in Milan. An appropriate weighting of the individual variances was applied in statistical analyses to correct for sex ratio unbalances. All groups participated on a voluntary basis and each individual gave a written informed consent. The experimental procedure was approved by the Ethical Committees of the Geneva University Hospitals and of the UHSR University.

Apparatus and task

Scribbling movements were recorded with two similar digitising tables (in Geneva: Model #2200-2436, Numonics; size: 110 × 80 cm, resolution: 0.0025 cm, sampling rate: 200 samples/s; in Milan: Model #9240, CalComp; size: 61 × 61 cm, resolution: 0.0025 cm, sampling rate: 200 samples/s) placed horizontally in front of the seating participant. The chair height was adjusted individually to maximize comfort. The writing implement resembled an ordinary ballpoint pen. The position of the pen's tip was sampled continuously as long as the tip was within a proximity margin of 13 mm. The experimenter placed successively at the centre of the workspace ten sheets of paper on which were printed circular frames of different sizes. The areas of the circles increased in a logarithmic progression from $\pi \text{ cm}^2$ (radius = 1 cm) to $81\pi \text{ cm}^2$ (radius = 9 cm). A different pseudo-random order of presentation was selected for each participant. Sequences complied with two constraints: (1) the first item should be neither the smallest, nor the largest circle; (2) the difference in rank order between successive items should be >2 .

The task was to draw a continuous smooth scribble inside each frame with the dominant hand. We instructed participants to make their drawings as meaningless as possible and to avoid repeating regular patterns. We also encouraged participants to avoid cusps in the trajectories. The movement was prompted by a “Go” verbal instruction, and went on until a “Stop” instruction. The recording lasted 10 s (2,000 samples). To eliminate the initial transient phase, the recording started a few seconds after the movement had actually begun. The entire experiment lasted about 15 min. Participants were free to choose the velocity at which the drawing was to be executed. As a consequence, average velocity varied substantially across individuals. In a few cases, participants inadvertently crossed the border of the limiting frame. These trials were repeated immediately. The experimenter paced the task by positioning the sheets with the traces. On average, there were 30 s between successive trials. At the beginning of the session, the task was introduced verbally by the experimenter who also demonstrated the functioning of the recording table. The actual recording session was preceded by few warm-up trials to familiarize the participants with the task and the material.

Groups PK and AMC repeated the task twice, in successive sessions, which also included two additional motor tasks (the results of which are not presented here). In session 1, which lasted about 75 min, PD patients were tested before taking their usual morning dose of levodopa and most, but not all, were in a variably severe OFF state corresponding to impaired motor function and obvious parkinsonism. First, they underwent a neuropsychological screening (see later). Then, they performed the three motor tasks. AMC participants followed the same routine. In session 2, PD patients were tested at least 90 min after taking their usual morning dose of levodopa (ON state). Clinical conditions in ON and OFF states were assessed by the Unified PD Rating Scale (UPDRS), part III (motor part, 27 items for a total score of 108 points) and the Purdue pegboard test with the left, right, and both hands. For patients the second session was an exact repeat of the first one. AMC participants also repeated the three motor tasks, but skipped the second neuropsychological screening. The two sessions were separated by a 30-min rest period. The notation PK1, PK2, AMC1, and AMC2 is used to indicate the results from the two sessions. The group of young controls (YC) performed the task only once and did not undergo a neuropsychological screening.

Neuropsychological test of the participants in groups PK and AMC

The neuropsychological screening was conducted by one of us (S.C.C.). It included a Stroop test and a test of verbal

Table 1 Population of age-matched controls and PD patients

AM controls: personal data				Pegboard			Stroop test				Verbal fluency
Name	Age	Sex	Height	Right	Left	Both	Points	Words	C. words	Errors	
af	61	F	165	12	11	9	15.00	23.00	34.00	0.00	1.57
cc	67	M	168	11	15	8	13.14	13.52	26.72	4.00	0.59
dm	82	F	175	13	11	8	12.34	31.41	49.17	0.00	−0.09
gf	68	M	176	13	12	9	11.30	24.38	29.02	0.00	−0.38
gk	58	M	180	13	11	10	12.91	16.00	26.12	2.00	1.70
gr	68	M	171	13	14	10	15.65	18.70	40.30	3.00	−2.02
ja	70	F	156	14	14	11	13.66	16.30	42.48	0.00	0.63
jo	67	M	182	12	10	8	18.06	28.53	57.45	4.00	−0.04
ld	80	M	172	10	12	9	13.82	17.67	20.53	0.00	1.26
mb	73	M	178	13	11	9	14.39	16.84	26.59	0.00	1.12
mo	76	M	173	13	12	11	12.61	19.63	28.91	1.00	0.10
pb	62	M	178	14	12	10	9.96	11.80	19.11	0.00	1.53
pc	57	M	164	14	16	12	12.30	19.31	27.83	1.00	−1.30
pu	57	M	175	13	12	8	13.06	13.62	23.10	0.00	−0.10
rl	64	F	158	13	10	9	14.57	17.50	25.92	0.00	0.90
ro	67	F	162	18	14	9	12.03	15.59	41.63	1.00	−1.92
sc	67	F	164	15	14	10	13.40	21.88	54.14	10.00	−1.26
si	72	M	170	14	11	10	13.41	17.04	24.91	0.00	−0.65
st	67	M	172	15	12	10	14.37	16.51	26.70	2.00	−1.62
sz	64	M	175	13	14	11	14.56	27.01	25.31	0.00	−2.17
Mean	66.9	14/6	171.05	13.10	12.05	9.05	13.52	19.46	32.05	1.27	−0.05

PD patients: personal data					Pegboard			Stroop test				Verbal fluency
Name	Age	Sex	Years	Height	Right	Left	Both	Points	Words	C.words	Errors	
bz	68	F	9	166	7	11	5	13.51	11.72	33.34	1.00	−1.28
ci	67	M	8	172	12	11	6	17.96	21.03	35.73	0.00	−1.13
ct	77	M	3	169	10	7	7	15.64	22.80	29.54	0.00	0.54
de	79	F	6	166	8	8	7	16.17	20.02	35.43	2.00	−2.39
dp	72	M	5	187	9	9	9	15.02	17.20	24.13	1.00	−0.03
dz	73	M	7	178	8	8	8	14.07	15.60	22.93	0.00	1.03
fk	61	F	17	175	7	9	6	12.03	14.72	22.50	0.00	0.74
gb	62	F	3	163	13	12	11	11.22	12.91	19.16	0.00	−0.42
hu	85	M	11	165	5	6	4	29.06	33.53	57.72	7.00	0.20
mm	62	M	5	178	9	9	7	15.75	21.40	30.96	1.00	−0.17
mq	71	M	8	169	6	7	5	34.11	32.13	34.06	4.00	−2.01
od	70	F	10	158	10	9	6	12.00	19.96	35.00	2.00	−1.00
ol	72	M	5	172	10	9	7	19.28	37.96	52.41	5.00	−0.88
on	62	M	11	169	12	10	7	10.84	16.50	30.55	3.00	−0.58
pe	57	F	2	158	14	11	9	15.84	14.41	17.75	1.00	−0.77
pf	68	M	14	178	6	5	2	25.63	37.41	61.53	6.00	−0.89
po	79	M	8	192	12	11	8	15.41	20.14	22.57	0.00	−1.16
pr	71	M	5	180	8	7	7	21.93	22.80	46.10	3.00	−1.30
tn	53	M	3	179	6	8	4	11.83	18.11	43.56	1.00	1.53
yc	59	F	24	163	12	10	8	13.24	13.15	24.47	1.00	0.40
Mean	68.29	13/7	8.20	172.1	9.25	9.10	6.15	17.03	21.17	33.91	1.89	−0.47

Personal data and scores for Pegboard, Stroop, and verbal fluency tests

fluency. Tables 1 and 2 summarize the results of the screening, as well as the UPDRS III scores and pegboard test results. The scores of the two groups in the Stroop test were similar (coloured points: PK = 17.03 s, AMC = 13.52 s; words: PK = 21.17 s; AMC = 19.46 s; coloured words: PK = 33.91 s; AMC = 32.05 s; errors with coloured words: PK = 1.89; AMC = 1.27). The only significant difference was the time for completing the coloured points test ($F(1,40) = 5.60$, $P = 0.0230$). Individual scores for the fluency test were ipsitized using the reference values for age, sex and education degree reported by Cardebat et al. (1990). Statistical analysis revealed no significant difference between groups (PK = -0.47; AMC = -0.05; $F(1,40) = 1.52$, $P = 0.22$). By contrast, the Purdue pegboard test scores, that correlate with dopaminergic deficit in the nigro-striatal system (Vingerhoets et al. 1997), were significantly different in the two groups (right hand: PK = 9.25; AMC = 13.10; $F(1,40) = 35.22$, $P < 0.0001$; left hand: PK = 9.10; AMC = 12.05; $F(1,40) = 42.36$, $P < 0.0001$; both hands: PK = 6.15; AMC = 9.05; $F(1,40) = 31.91$, $P < 0.0001$). The UPDRS scores clearly discriminated the age-matched controls from the patients in both ON and OFF states (AMC: range = 0–5.5; mean = 2.7. PK-OFF state: range = 7–43.5; mean =

23.1; PK-ON state: range = 5–37; mean = 17.8). The difference between scores in ON and OFF states was highly significant ($F(1,40) = 86.14$, $P < 0.0001$). However, the reduction of the UPDRS scores in the ON state varied considerably among patients (range 3–64%). The sub-scores of the UPDRS indicated in Table 2 also differed between groups PK and AMC. Axial bradykinesia was computed according to Stebbins and Goetz (1998) as the sum of items 18 (speech), 19 (facial expression) and 27–31 (arise from chair, posture, gait, postural stability, body bradykinesia). Right bradykinesia was the sum of items 23r, 24r, 25r, and 26r (finger taps, hand grip, hand pronate/supinate and leg agility). Rigidity, rest tremor and action tremor were computed from items 22, 20 and 21, respectively. In all cases, except for rest tremor (PK = 1.01; AMC = 0.00; $F(1,40) = 6.03$, $P = 0.0185$) and action tremor (PK = 0.75; AMC = 0.05; $F(1,40) = 4.17$, $P = 0.0477$), differences were highly significant, at the 0.0001 level.

Data processing

The data from each trial (2,000 pairs of coordinates of the point-samples) were processed off-line. The parameters

Table 2 Population of PD patients

Name	UPDRS		Axial brady		Right brady		Left brady		Rigidity		Rest tremor		Action tremor	
	1	2	1	2	1	2	1	2	1	2	1	2	1	2
bz	15.0	11.5	4.0	4.0	1.0	1.0	4.0	1.0	5.0	4.5	1.0	1.0	0.0	0.0
ci	18.0	15.0	1.0	1.0	4.5	3.0	3.5	3.0	5.0	4.0	4.0	4.0	0.0	0.0
ct	24.5	22.5	8.5	8.5	3.0	3.0	5.0	4.5	8.0	6.5	0.0	0.0	0.0	0.0
de	11.5	9.5	4.0	4.0	2.0	1.5	3.5	1.5	2.0	2.5	0.0	0.0	0.0	0.0
dp	16.0	14.5	4.0	4.0	2.0	2.0	2.0	2.5	7.5	6.0	0.0	0.0	0.5	0.0
dz	27.5	20.0	6.0	5.0	3.0	1.0	7.0	4.5	3.5	3.5	8.0	6.0	0.0	0.0
fk	43.5	21.5	8.5	5.5	11.0	7.0	2.5	1.0	11.5	5.5	3.5	1.0	6.5	1.5
gb	11.0	4.0	1.5	0.0	3.5	3.0	0.0	0.0	4.5	1.0	1.5	0.0	0.0	0.0
hu	26.5	17.0	12.5	8.0	4.0	3.0	3.0	2.5	5.0	3.5	0.0	0.0	2.0	0.0
mm	24.5	18.5	6.5	5.5	4.0	3.0	8.0	6.0	6.0	4.0	0.0	0.0	0.0	0.0
mq	24.0	20.0	5.0	5.0	4.0	3.0	7.0	5.0	8.0	7.0	0.0	0.0	0.0	0.0
od	23.0	17.0	8.0	7.0	4.0	3.0	6.5	4.5	3.5	2.5	1.0	0.0	0.0	0.0
ol	28.5	26.5	11.0	11.0	6.0	5.0	6.5	5.0	5.0	5.5	0.0	0.0	0.0	0.0
on	7.0	5.5	2.5	2.0	0.0	0.0	2.5	1.5	2.0	2.0	0.0	0.0	0.0	0.0
pe	9.0	5.0	0.5	0.0	0.0	0.0	3.0	2.5	4.5	2.0	1.0	0.5	0.0	0.0
pf	38.0	37.0	6.0	6.5	12.0	11.5	10.0	10.5	8.0	8.5	0.0	0.0	2.0	0.0
po	31.5	23.0	5.0	5.0	5.5	5.0	7.0	4.0	12.0	7.0	1.0	1.0	1.0	1.0
pr	21.5	20.0	6.0	6.0	6.0	5.0	5.5	5.0	3.0	3.0	0.0	0.0	1.0	1.0
tn	40.5	37.0	9.5	9.5	13.5	13.0	10.0	8.0	5.5	5.5	1.0	0.0	1.0	1.0
yc	21.0	11.0	6.5	3.5	7.0	4.0	3.5	1.5	3.0	1.5	0.0	0.0	1.0	0.5
Mean	23.1	17.8	5.77	5.00	4.79	3.80	5.05	3.70	5.60	4.30	1.01	0.70	0.75	0.20

Personal data and UPDRS scores

selected for characterizing the scribbling movements were: (1) the instantaneous tangential velocity $V(t)$; (2) the number of inflections in the trace N_f ; (3) the length L_{seg} of the segments of trajectory bounded by successive inflection points; (4) the normalized radius of curvature of the trace $R^{*1/3}$; (5) the average curvature of the trace C_m ; (6) the average velocity gain factor over the entire trace (K_m) and over successive segments (K_{seg}). Appendix 1 provides a detailed definition of these parameters and a description of the data processing for their estimation.

Results

All participants found the experimental task easy to understand and to execute. In young participants, very few trials had to be repeated, either because the trace had crossed the circular frame, or because the smoothness condition (i.e. no cusps in the trajectory) had been violated. The number of repeated trials (mostly because of border crossings) was somewhat higher for both age-matched controls and PD patients. The database for each group and

session (YC, AMC1, AMC2, PK1, PK2) was a set of 20 (subjects) \times 10 (trials) = 200 records. Figure 1 shows the complete set of traces produced by a typical young control. Figure 2 shows the corresponding results for one patient. This figure exemplifies the typical behaviour of PD patients in the OFF state, namely slower, more hesitating movements with a higher number of small direction changes. Signs of micrographia and stereotyped, repetitive patterns were also common in the traces of patients in both ON and OFF states.

Figural self-similarity

After scaling the traces to the same size, visual inspection of individual performances suggested that the shape of the movements was fairly invariant across frame sizes (self-similarity). As shown for one typical individual in group YC (upper panels in Fig. 3), and session 2 of group PK (upper panels in Fig. 4) there was no obvious figural clue to tell which drawing was made within the largest or the smallest frame. Individual differences in young and age-matched controls were also small. This is exemplified by the traces in the lower panels of Fig. 3 generated by two young participants within the same frame. The lower

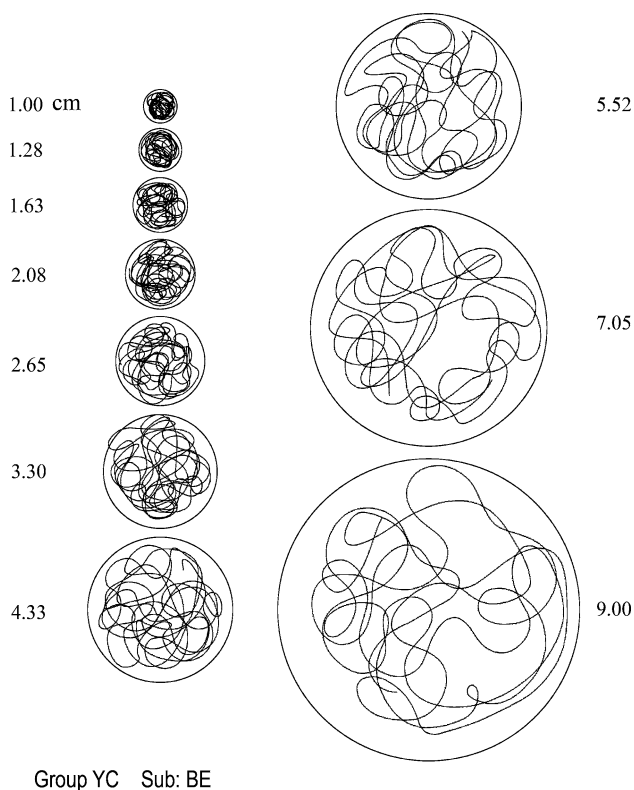


Fig. 1 Scribbling movements. A complete series of recordings in one representative young control. The radius of the circular frames increased logarithmically from 1 to 9 cm. During the experiment only one frame at the time was presented on the recording tablet. The sequence with which different sizes were presented was varied randomly for each participant

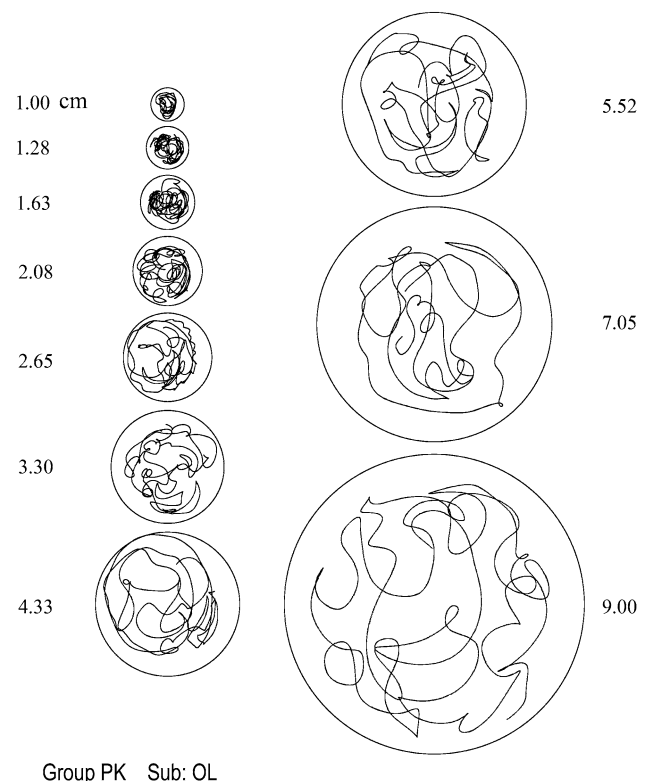


Fig. 2 Scribbling movements. A complete series of recordings in one representative PD patient in the OFF state. Traces in patients were far less fluent than in controls

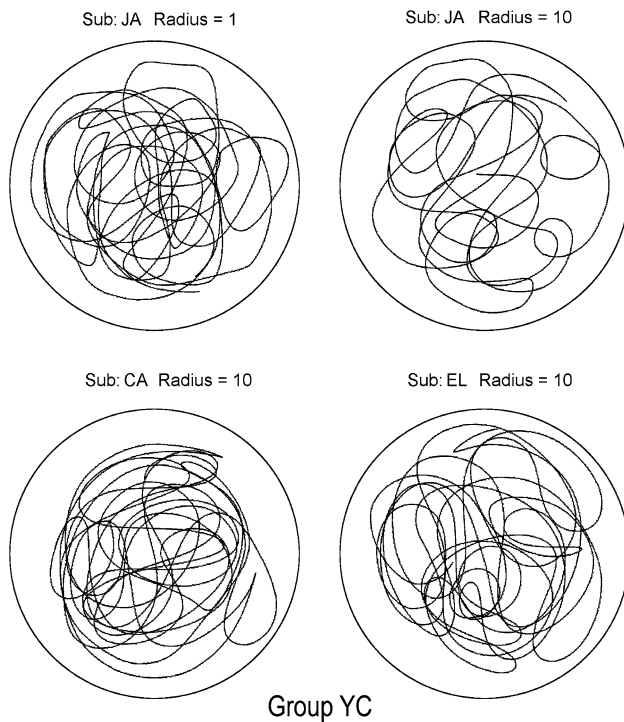


Fig. 3 Figural similarity. *Top traces* example in one representative young control (JA) of the self-similarity of traces at the extremes of the size range. For clarity, the two frames have been scaled to the same size. *Bottom traces*: scribbles from two young controls (CA and EL) illustrate the range of inter-individual variability in normal participants

panels in Fig. 4 illustrate the inter-individual similarity observed in the PK group (see, however, above). Figural self-similarity across sizes was assessed quantitatively in two ways.

First, we explored how the number of inflections N_f varied as a function of frame size. Because movements were constrained by a frame, this parameter depends on the general shape of the scribbles. Few inflections imply the presence of many loops in the trace, whereas a high number of inflections suggest that the trajectory was wavy and sinuous. A preliminary analysis of the data showed that all movement parameters varied approximately as a power function $Y = K \times A^\gamma$ of the area A of the frame. Thus, for this and all subsequent analyses we estimated the parameters K and γ by computing the linear regression $\log(Y) = a + b \log(A)$ and setting $K = e^a$, $\gamma = b$. The average and between-participant variability of N_f was considerably higher in AMC and PK groups than in group YC. Yet, for all groups the power function approximation predicted well the dependence of N_f on the frame area (YC: $N_f = 23.742 A^{-0.069}$; AMC1: $N_f = 47.790 A^{-0.120}$; AMC2: $N_f = 48.671 A^{-0.130}$; PK1: $N_f = 48.200 A^{-0.149}$; PK2: $N_f = 37.863 A^{-0.149}$). In all cases the exponent of the power law was small, indicating that the number of inflections was weakly dependent on movement size.

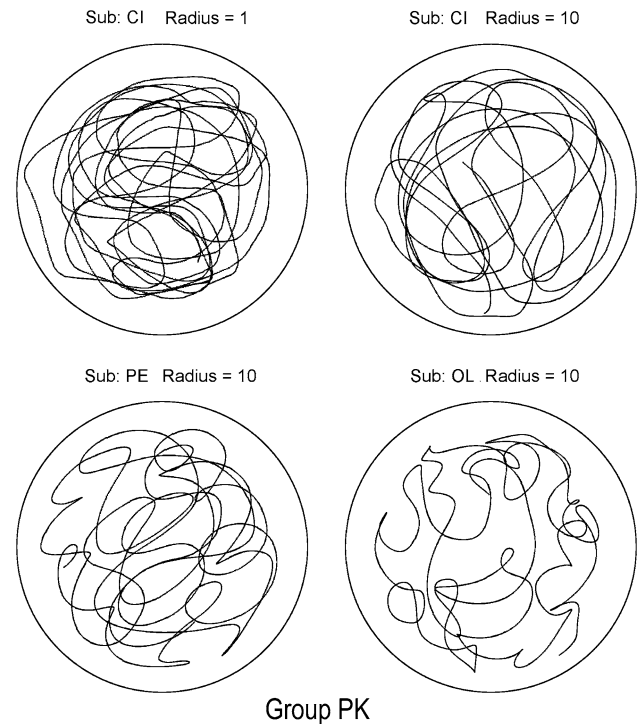
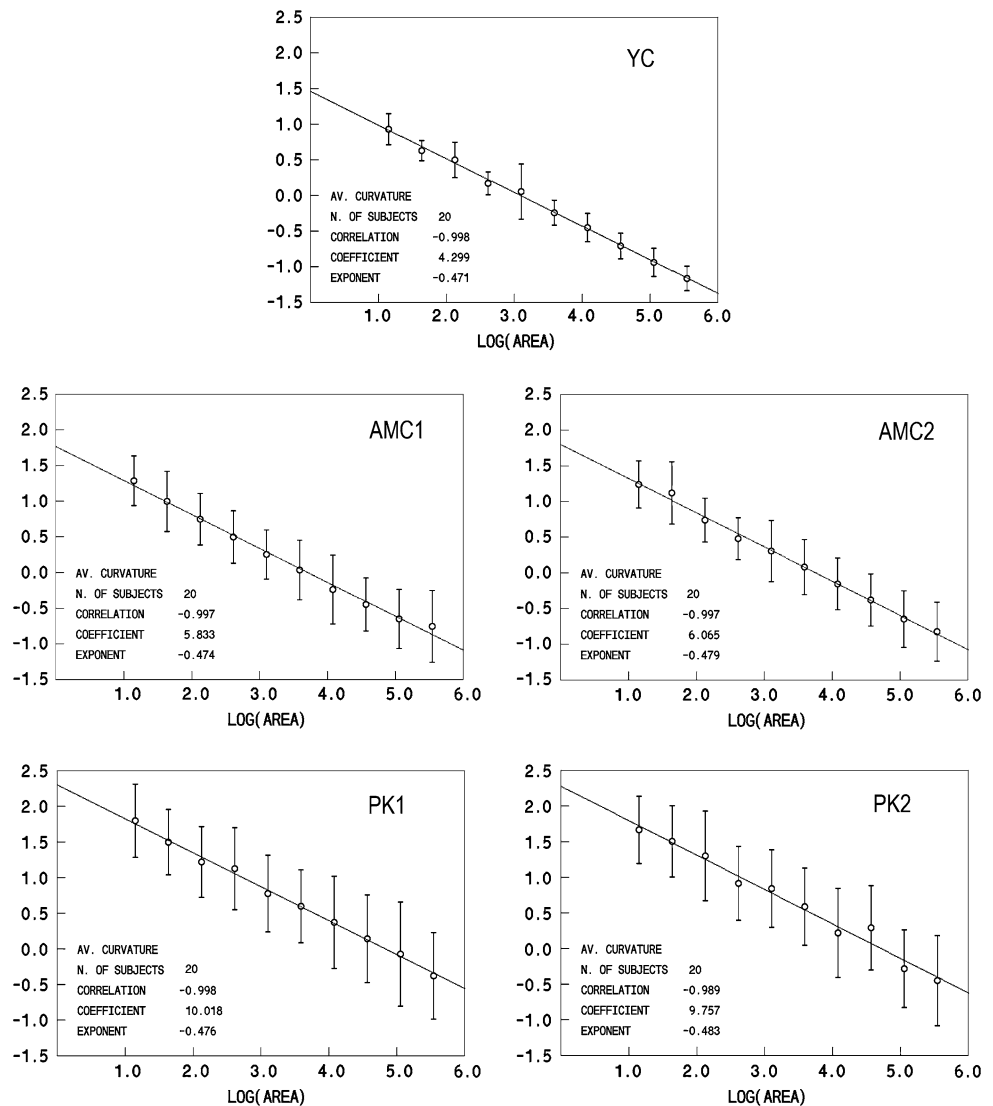


Fig. 4 Figural similarity. *Top traces* example in one representative patient (CI) of the self-similarity of traces at the extremes of the size range. For clarity, the two frames have been scaled to the same size. *Bottom traces* scribbles from two patients in the OFF state (PE and OL) illustrate the range of inter-individual variability in patients

The second figural index for assessing self-similarity was the average curvature of the traces C_m computed with the re-sampling technique detailed in Appendix 1. Figure 5 shows for each group the relation between $\log(C_m)$ and $\log(A)$ (C_m values averaged over participants). The excellent fit of the linear regression demonstrates that also in this case a power function predicts accurately the average curvature for each frame size (YC: $C_m = 4.299 A^{-0.471}$; AMC1: $C_m = 5.833 A^{-0.474}$; AMC2: $C_m = 6.065 A^{-0.479}$; PK1: $C_m = 10.018 A^{-0.476}$; PK2: $C_m = 9.757 A^{-0.483}$). C_m can also be expressed as a power function of the (constant) curvature of the frame (C_f) by using the relation $A = \pi C_f^{-2}$ (YC: $C_m = 2.507 C_f^{0.942}$; AMC1: $C_m = 3.390 C_f^{0.948}$; AMC2: $C_m = 3.505 C_f^{0.958}$; PK1: $C_m = 5.809 C_f^{0.952}$; PK2: $C_m = 5.613 C_f^{0.966}$). The average curvature was significantly different among groups, young subjects and PD patients producing the lowest and highest curvature, respectively. However, the fact that the exponent of the ($C_m - C_f$) relationship was always close to 1 demonstrates that all participants spontaneously scaled almost linearly the average curvature of their movements to the curvature of the frame. Insofar as the curvature is concerned, scribbles of different dimensions were just scaled versions of an idiosyncratic basic pattern. Across participants, the variability of C_m (estimated by the 95% confidence intervals

Fig. 5 Relation between the logarithm of frame area and the logarithm of the average curvature for the three groups of participants. For groups AMC and PK the results of the two sessions are shown separately. Data points are averages over all participants. Bars encompass ± 1 SD. In all cases a linear regression fits very accurately the data points. The parameters of the power law corresponding to the fit in linear scales are indicated inset. Note that only the coefficient, but not the exponent of the power law differs significantly among groups



around the means) was surprisingly small for the group YC, and moderate for the other groups, indicating a reasonable degree of homogeneity of the individual performances. Similar results were also observed for N_f . Thus, we felt confident that the group results to be presented in the next sections capture some common features of the motor behaviour under study.

Scribble comply with the Two-third Power Law

The velocity of end-point voluntary movements is modulated both globally, by setting the average velocity for the entire gesture, and locally, by the geometrical properties of the trajectory (see “Introduction”). The emphasis in this study is on the former type of control. However, the two modulations are intertwined. To isolate the control component acting on average velocity, the first preliminary step

was to verify in both patients and controls that the Two-Third Power Law $V(t) = K[R(t)/(1 + \alpha R(t))]^{(1-\beta)}$ applies also to scribbling movements. The law states that movement velocity is the product of a *velocity gain factor* K , which sets the average velocity, and of a time-varying term describing the instantaneous modulation that depends on the radius of curvature of the trajectory $R(t)$. The parameter α in the equation above has a technical role. Its introduction was suggested by Viviani and Stucchi (1992) to generalize the original formulation of the Power Law $V(t) = K \times R(t)^{1-\beta}$ also to trajectories with points of inflection where the radius of curvature has a singularity. Instead, the expression $R^*(t) = [R(t)/(1 + \alpha R(t))]$ stays finite for every value of $R(t)$. Viviani and Stucchi (1992) showed that law is most accurate when the parameter α is allowed to depend on the average velocity V_m . To assess the validity of the law, we considered α , β , and K in the equation above as

free parameters. Using a Simplex algorithm, we estimated for each trial the values of the parameters that minimize the mean square difference between actual and predicted velocities. Figure 6 illustrates with an example from a young control the regression between $\log V(t)$ and $\log([R(t)/(1 + \alpha R(t))])$ after inserting the optimal value of α . In this diagram the slope of the regression line estimates β and the intercept at the origin estimates $\log(K)$.

As expected, the optimal estimate of α varied as a function of frame size (ANOVA, YC: $F(9,171) = 38.57$, $P < 0.001$; AMC1: $F(9,171) = 41.91$, $P < 0.001$; AMC2: $F(9,171) = 22.40$, $P < 0.001$; PK1: $F(9,171) = 18.94$, $P < 0.001$; PK2: $F(9,171) = 55.37$, $P < 0.001$). In addition, there were significant differences among participants (ANOVA, YC: $F(19,171) = 5.31$, $P < 0.001$; AMC1: $F(19,171) = 11.65$, $P < 0.001$; AMC2: $F(19,171) = 6.60$, $P < 0.001$; PK1: $F(19,171) = 8.81$, $P < 0.001$; PK2: $F(19,171) = 10.32$, $P < 0.001$). For all trials and all participants in each group the logarithms of optimal α -values were linearly regressed against the logarithms of the corresponding average velocities (we excluded from the analysis a few trials in which $\alpha > 2.5$ because inspection of the data showed that the corresponding movements were abnormally slow). In spite of individual differences (see above), the power function approximation that in linear scales corresponds to the logarithmic regression was remarkably similar for all groups (YC: $\alpha = 6.315 V_m^{-1.007}$; AMC1: $\alpha = 8.793 V_m^{-1.038}$; AMC2: $\alpha = 8.602 V_m^{-1.046}$; PK1: $\alpha = 5.403 V_m^{-1.027}$; PK2: $\alpha = 6.539 V_m^{-1.061}$). In fact,

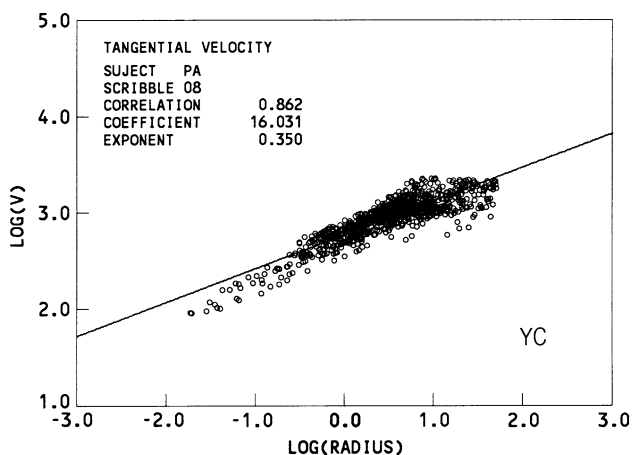
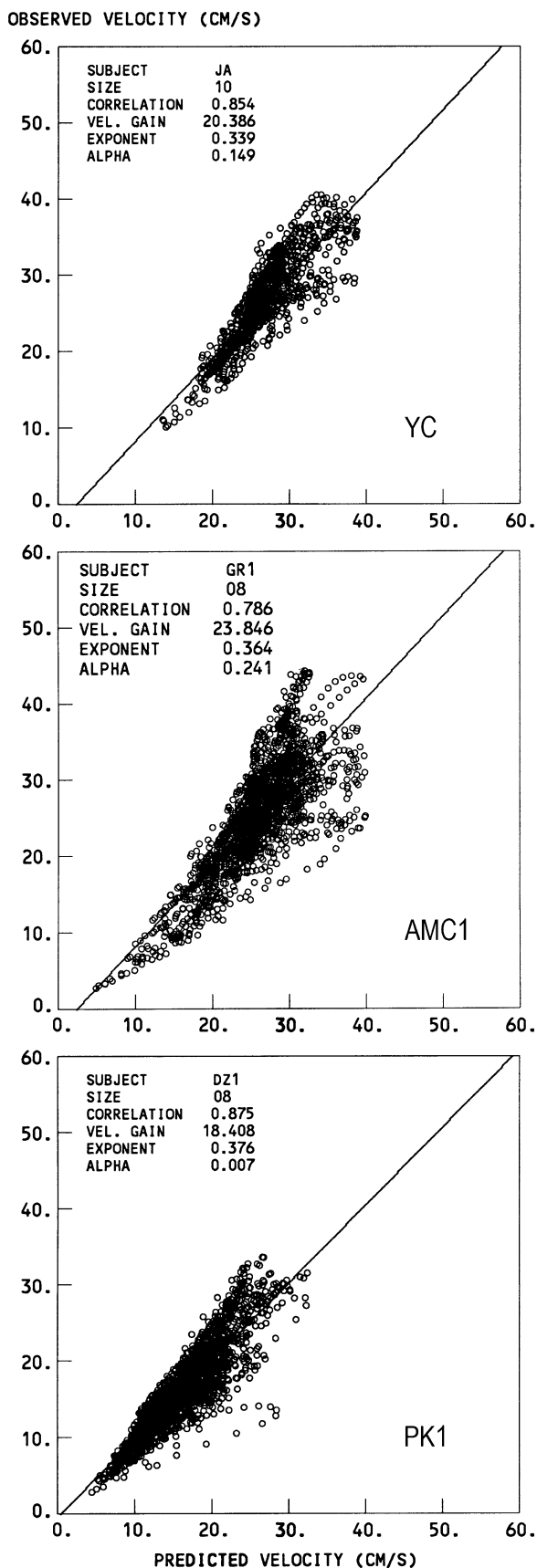


Fig. 6 Scribbles comply with the Two-thirds Power Law. Data from one representative participant (group YC) and for the indicated frame size. Scatter diagram (2,000 points) between the logarithm of the instantaneous velocity (V ordinate) and the logarithm of the generalized radius of curvature (R abscissa). A linear regression fits very accurately the data points. The parameters of the power law corresponding to the fit in linear scales are indicated inset. Note that the exponent $1 - \beta$ of the power law is quite close to the theoretical value $1/3$

the values of α averaged over frame size (YC: 0.796; AMC1: 0.904; AMC2: 0.910; PK1: 1.001; PK2: 0.975) were not significantly different across groups [one-way ANOVA, $F(4,95) = 0.589$, $P = 0.671$]. The introduction of this parameter was instrumental to keep the Power Law well behaved near the points of inflection. Also, letting it vary across frame sizes and groups improved the fit of the Power Law. However, keeping it constant for all trials in all participants would not have obscured the significant modulation of the velocity gain among groups (see below).

As for the exponent β , there was a slight albeit systematic non-linear effect of the frame size (ANOVA, YC: $F(9,171) = 28.60$; $P < 0.001$; AMC1: $F(9,171) = 15.68$, $P < 0.001$; AMC2: $F(9,171) = 21.72$, $P < 0.001$; PK1: $F(9,171) = 6.69$, $P < 0.001$; PK2: $F(9,171) = 16.54$, $P < 0.001$), as well as significant individual differences (ANOVA, YC: $F(19,171) = 4.65$, $P < 0.001$; AMC1: $F(19,171) = 8.70$, $P < 0.001$; AMC2: $F(19,171) = 10.81$, $P < 0.001$; PK1: $F(19,171) = 5.98$, $P < 0.001$; PK2: $F(19,171) = 6.54$, $P < 0.001$). However, the exponent of the Two-thirds Power Law was virtually indistinguishable across groups [YC: $1 - \beta = 0.3526$; AMC1: $1 - \beta = 0.3572$; AMC2: $1 - \beta = 0.3576$; PK1: $1 - \beta = 0.3527$; PK2: $1 - \beta = 0.3552$; ANOVA, $F(4,95) = 1.101$, $P = 0.361$], and very close to the theoretical value $\beta = 2/3$. More importantly, for all groups the variability was no more than 3% of the corresponding average. Finally, significant differences among groups emerged in the least-square estimate of the velocity gain factor K (YC: $K = 13.723$; AMC1: $K = 16.724$; AMC2: $K = 16.632$; PK1: $K = 12.089$; PK2: $K = 12.809$; ANOVA, $F(4,95) = 4.821$, $P = 0.001$). These differences will be analysed further in the next section. The accuracy with which the Two-thirds Power Law accounted for the data was estimated by regressing observed against predicted velocity. The coefficient of linear correlation was uniformly high for all trials (YC: $r = 0.671 \pm 0.107$; AMC1: $r = 0.674 \pm 0.080$; AMC2: $r = 0.672 \pm 0.078$; PK1: $r = 0.703 \pm 0.086$; PK2: $r = 0.695 \pm 0.085$). Figure 7 illustrates these regressions with one example from groups YC, AMC1, and PK1.

In summary, even for highly variable and extemporaneous scribbling movements, the instantaneous velocity is related to the corresponding value of the radius of curvature as prescribed by the Two-thirds Power Law. PD patients (group PK1 and PK2) had a consistently lower gain than both age-matched and young controls, but the relational constraint described by the law appears to be robust against age and clinical conditions. This result provides the basis for the timing analysis to be presented next.



◀ **Fig. 7** Scribbles comply with the Two-thirds Power Law. Data from one representative participant for each group and for the indicated frame sizes. Scatter diagram (2,000 points) between the observed instantaneous velocity (ordinates) and the instantaneous velocity predicted by the Two-thirds Power Law (abscissas). In all cases a linear regression with a slope very close to 1 fits very accurately the data points. The parameters of the power law corresponding used to compute the predicted velocity are indicated inset. Note that the exponents $1 - \beta$ of the power law are all close to the theoretical value $1/3$

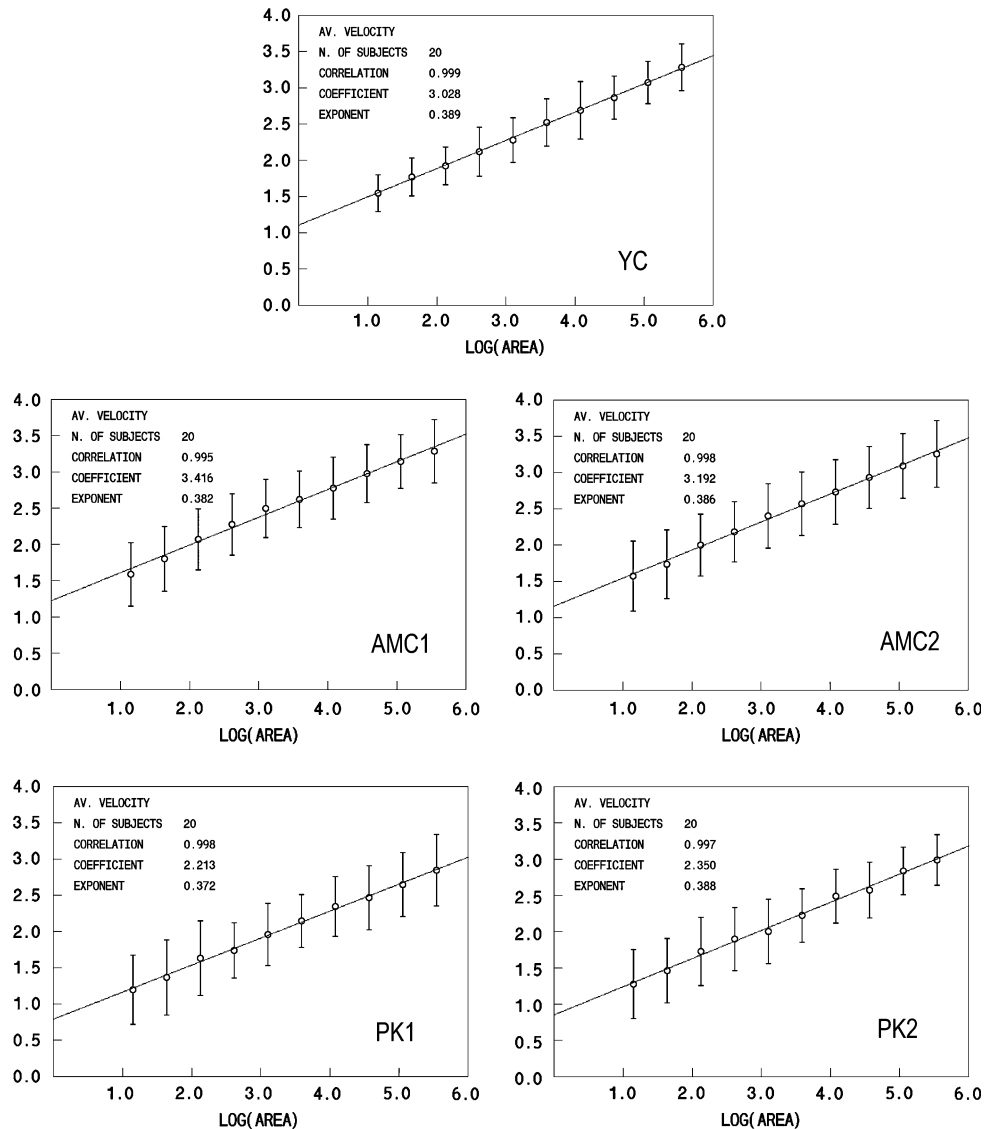
Global isochrony

For point-to-point movements the phenomenon of isochrony manifests itself as a compensatory increase of the average velocity with the length of the trajectory to be executed. In the case of scribbles whose trajectory is not defined in advance we have to envisage the possibility that the role of path length for velocity scaling is taken by a more general figural parameter that can be represented centrally. We investigated the hypothesis that the frame surface is such a parameter by testing whether the surface permits one to predict reliably the average velocity V_m . As before, the test involved computing the linear regression between $\log(A)$ and $\log(V_m)$. Figure 8 shows the results for all experimental groups. The coefficient of linear correlation for each data set was very high (see inset). Thus, the power functions approximations derived from the regression (YC: $V_m = 3.028 A^{0.389}$; AMC1: $V_m = 3.416 A^{0.382}$; AMC2: $V_m = 3.192 A^{0.386}$; PK1: $V_m = 2.213 A^{0.372}$; PK2: $V_m = 2.350 A^{0.388}$) are quite accurate. The multiplicative coefficients in these approximations reflect faithfully the differences among the group estimates of the velocity gain factor K in the Two-thirds Power Law (see previous section). By contrast, the exponents were almost identical across groups.

The same analysis was also carried out for each participant separately. The results demonstrated that the velocity compensation emerging at the group level was in fact present in each individual performance. The multiplicative velocity gain factor was somewhat variable across individuals because each of them selected a different overall tempo for the task. There were small variations also in the exponent of the power function, suggesting that the degree of velocity compensation was not uniform within the population. However, the standard deviation of the exponent was far smaller than that of the multiplicative factor. In other words, individual linear regressions were almost parallel to each other. More importantly, the standard deviation of the coefficient of correlation was quite small, and the average of the individual correlations was not much lower than the correlation for the group data. Clearly, the high regularity of the results in Fig. 8 is not an

Fig. 8 Global isochrony.

Relation between the logarithm of the frame area (abscissas) and the logarithm of the average velocity. For groups AMC and PK the results of the two sessions are shown separately. Data points are averages over all participants within a group. Bars encompass ± 1 SD. In all cases a linear regression fits very accurately the data points. The parameters of the power law corresponding to the fit in linear scales are indicated in inset. Note that the exponent of the power law is similar for all groups. Averaged over trials and frame sizes, the velocity was YC: 12.84 cm/s; AMC1: 14.08 cm/s; AMC2: 13.37 cm/s; PK1: 8.76 cm/s; PK2: 9.92 cm/s



artifactual consequence of pooling individual data. In summary, the analysis confirmed the hypothesis that the area of the frame affords a stable figural parameter that participants used to modulate the base rhythm in an (unconscious) effort to increase the velocity *pari passu* with the overall size of the movement. In particular, with respect to PD patients, these results suggest that the disease does not affect the mechanisms that modulate globally the velocity as a function of the available workspace. Next, we turn to the issue of whether this velocity compensation mechanism can be factored out in a manner similar to that demonstrated for closed, regular trajectories.

Decomposing the velocity compensation mechanism

By integrating the two sides of the Two-Thirds Power Law one gets

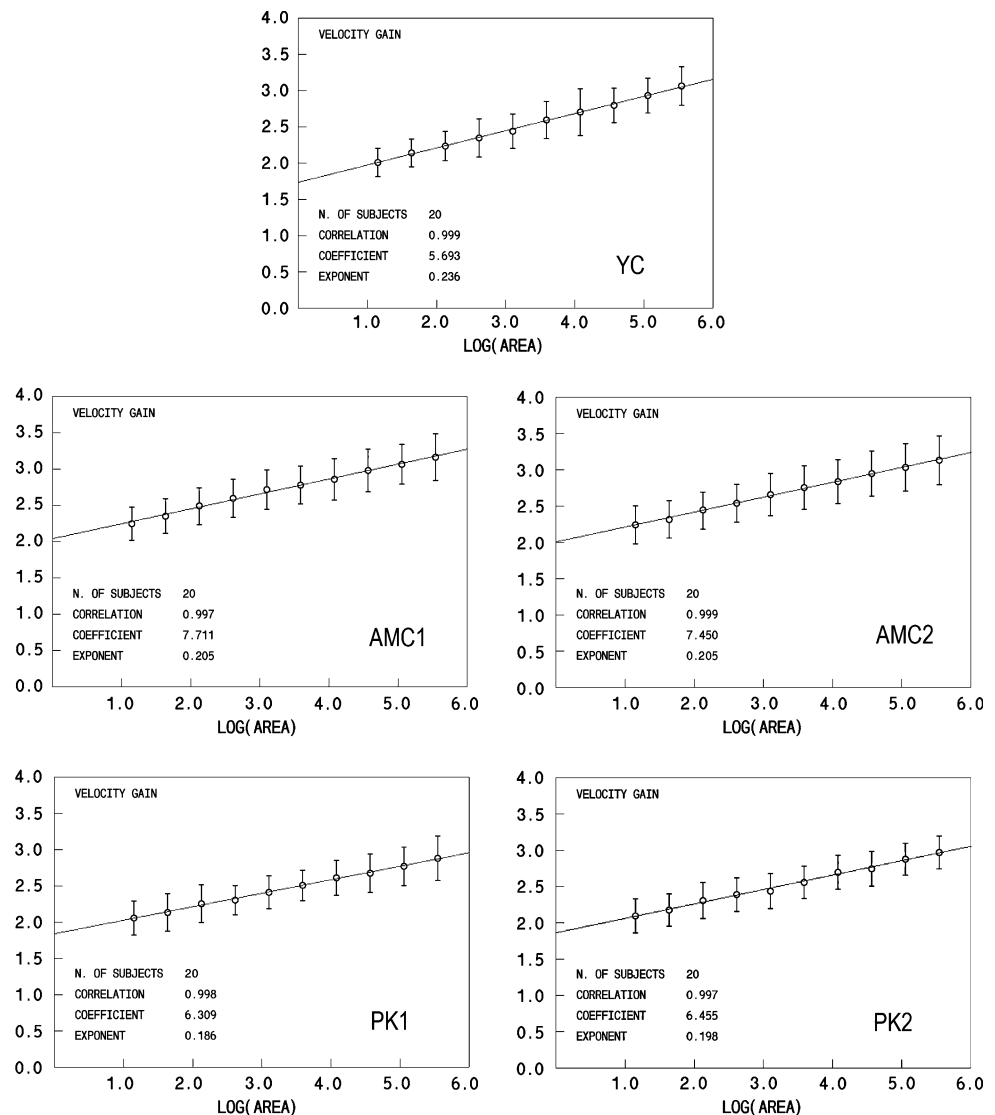
$$V_m = \frac{1}{T} \int_0^T V(t) dt = \frac{K}{T} \int_0^T R(t)^{1/3} dt = K \langle R(t)^{1/3} \rangle$$

showing that the average velocity can be decomposed in two multiplicative factors. In the case of closed regular trajectories, the increase of the average velocity with increasing size of the figure was found to result from the contributions of both factors. On the one side, the velocity gain factor K increases with trace length, which scales linearly with figure size. On the other side, increasing the size shifts upward the range of values of the radius of curvature, and, therefore, increases the value of the time average $\langle R^{1/3}(t) \rangle$. In past work (see “Introduction”) we argued that the contribution of the factor K reflects compensatory mechanisms acting at the motor planning stage, whereas the contribution of the radius of curvature emerges at the implementation stage. Thus, we reserved the term

“Isochrony” to describe only the effect of the gain factor K on average velocity. Here we verified that the distinction suggested by the analysis of closed regular trajectories remained valid also in the case of scribbles. Because the average curvature of the traces was inversely related to the frame area (see Fig. 5), the time average $\langle R^{*1/3}(t) \rangle$ is an equally regular increasing function of the frame area (recall that the curvature is the inverse of the radius). It follows that, even in scribbles, the factor $\langle R^{*1/3}(t) \rangle$ contributes to the increase of average velocity with frame area. To demonstrate that the factor K also provides its own distinct contribution, for each trial we computed directly the average velocity gain factor K_m by integrating the expression $V(t)/[(1 + \alpha R(t))/R(t)]^{1/3}$ over the recording interval $[0 - T]$. Then, we regressed the mean K_m across participants against the area A , after applying the usual logarithmic transformation of the variables. The results for all groups are illustrated in Fig. 9, which reports the

coefficients of the best-fitting power function interpolation to the data points: [YC: $K_m = 5.693 A^{0.236}$; AMC1: $K_m = 7.711 A^{0.205}$; AMC2: $K_m = 7.450 A^{0.205}$; PK1: $K_m = 6.309 A^{0.186}$; PK2: $K_m = 6.455 A^{0.198}$]. In all cases, the accuracy of the interpolation was excellent and the exponents of the power law were similar. Moreover, although we considered frame area rather than path length as a control variable, the results are dimensionally congruent with those reported previously for closed, periodic movements. For instance, in the case of ellipses (Lacquaniti et al. 1984), the relation between velocity gain factor K_m and perimeter P is well approximated by $K = K_0 \times P^{0.45}$. For eccentricities up to 0.9 the perimeter of the ellipse is almost proportional to the square root of its area. Thus, this relation can also be written as $K = K_0 \times A^{0.225}$, where the value of the exponent is in excellent agreement with the values estimated above. The agreement supports the suggestion made above that the role

Fig. 9 Decomposing the velocity compensation mechanism. Relation between the logarithm of the frame area (abscissas) and the logarithm of velocity gain factor. For groups AMC and PK the results of the two sessions are shown separately. Data points are averages over all participants within a group. Bars encompass ± 1 SD. In all cases a linear regression fits very accurately the data points. The parameters of the power law corresponding to the fit in linear scales are indicated inset. Note that the exponent of the power law differs significantly among groups. Averaged over trials and frame sizes, the velocity gain was YC: 13.22; AMC1: 15.94; AMC2: 15.40; PK1: 12.15; PK2: 12.99



of the control variable in scribbling is taken by the total surface within which the movement is constrained. If so, the expression $K_m = K_0 \times A^e$ would capture the isochronic component of the velocity modulation.

The velocity gain factor is independent of the average curvature

We showed that the frame area A has a graded effect both on the average radius of curvature of the traces $\langle R^{*1/3} \rangle$ and on the velocity gain factor K . However, in order to generalize also to scribbles the notion that K and $\langle R^{*1/3} \rangle$ reflect the working of different motor mechanisms (see above), one must show that the two factors are independent. This check was performed on the data of the group YC by an analysis of the residuals. First, we computed separately the regressions between $\langle R^{*1/3} \rangle$ and A , and between K_m and A , by taking into account the entire set of 20 (participants) \times 10 (frame sizes) = 200 data points. Then, we computed the regression between the residuals with respect to the corresponding regression lines. As shown in Fig. 10, there was no significant correlation between the residuals. As we shall see in the next section, the velocity gain factor is not constant within a trial. However, the lack of correlation shows that, globally, the gain was independent of the curvature of the trace. This conclusion provides further strong support to the contention that velocity is indeed the result of the multiplicative effect of the two factors K and $\langle R^{*1/3} \rangle$, and that the decomposition of the velocity suggested by the Two-thirds

Power Law captures a significant feature of the underlying motor plan.

A local form of Isochrony

In this section, we demonstrate that the velocity gain factor K is modulated in a principled fashion in the course of the movement. The motivation for this further analysis is the following. Presumably, long and complex movements such as writing are represented in motor memory as a collection of figural units integrated at the time of execution. It has been suggested that in this case velocity is modulated jointly by the total length of the trace and, at a more local level, by the length of each successive figural unit (Viviani 1986). In scribbling the figural units are selected randomly from the available motor repertoire rather than following a predetermined scheme. However, we hypothesised that in addition to the global control of velocity dictated by the frame that constrains the movement, a similar form of local isochrony is present also in scribbles.

Testing this intuition is not straightforward because there is no principled criterion for identifying the endpoints of figural units. Even when a geometrical feature strongly suggests a natural segmentation in two units (e.g. in the case of a double ellipse), the analysis showed (Viviani and Cenzato 1985) that the influence of path length on motor planning might extend over both units. To overcome this difficulty, we adopted the heuristic strategy that consists in assuming that each segment of trajectory comprised between two points of inflection is a figural unit. This somewhat arbitrary criterion makes sense, however, from the point of view of motor control because inflections mark the transition between opposite directions of rotation.

The analysis of local isochrony was carried out on the pooled results of all participants within a group (Table 3). For each frame size, we began by identifying the trace segments comprised between points of inflection. Segments that were either too short (less than 40% of the average length), or too long (more than 200% of the average length) were discarded. For each segment, we computed its length (L_{seg}) by integrating the original velocity of the trace, and its average gain factor (K_{seg}) from the re-sampled trace (re-sampling was again necessary to prevent an over-representation of the high-curvature samples). In keeping with the large difference between the numbers of points of inflection (see above), the average segment length was much longer in young controls than in patients, with the AMC group being somewhat intermediate. The difference among groups of the average segment gain reflected the difference in velocity documented in Fig. 8. In the second step of the analysis, we correlated segment length and segment gain across frame areas. Because the total trace length increased with the frame

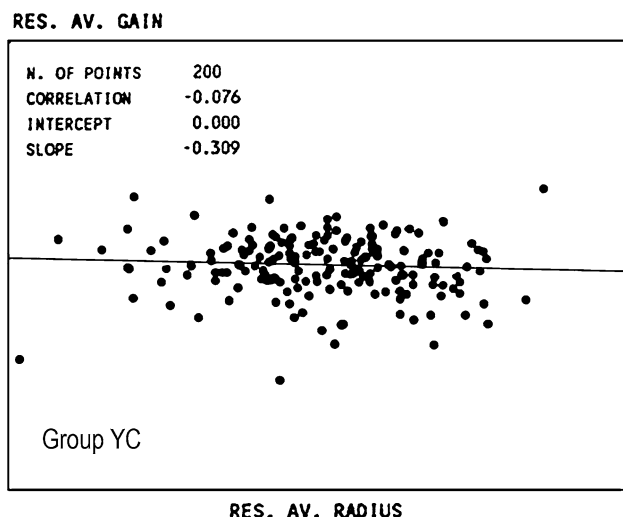


Fig. 10 Velocity gain factor is independent of average curvature. Scatter diagram of the residuals of the regression between $\langle R^{*1/3} \rangle$ and the frame area (abscissa) and the regression between K_m and the frame area (ordinate). Data for all participants in group YC. The residuals are not correlated

Table 3 Summary of segment analysis for all groups of participants

Group	YC	AMC1	AMC2	PK1	PK2
N	2,985	2,986	2,891	3,776	2,957
AL_{av}	7.212	4.678	5.023	1.504	2.014
AK_{seg}	13.11	16.33	16.10	9.90	10.98
$(K_{seg}/K_{av}) = c_0(L_{seg}/L_{av})^{\xi}$					
ρ	0.466	0.344	0.325	0.313	0.288
T_{max}/T_{min}	3.613	2.608	2.492	2.020	1.980
C_0	0.999	1.009	1.010	1.027	1.027
ξ	0.162 ± 0.0134	0.169 ± 0.0185	0.169 ± 0.0197	0.227 ± 0.0216	0.213 ± 0.0248
$K_{av} = c_1 A^{\theta}$					
ρ	0.813	0.704	0.733	0.694	0.736
C_1	5.160 ± 0.1608	5.027 ± 0.2285	4.813 ± 0.2193	3.388 ± 0.1298	2.992 ± 0.1297
θ	0.258 ± 0.0087	0.279 ± 0.0133	0.295 ± 0.0131	0.262 ± 0.0114	0.300 ± 0.0131
$L_{av} = c_2 A^{\psi}$					
ρ	0.724	0.680	0.674	0.616	0.669
C_2	1.245 ± 0.0853	0.327 ± 0.0355	0.401 ± 0.0427	0.148 ± 0.0137	0.103 ± 0.0119
ψ	0.428 ± 0.0192	0.632 ± 0.0317	0.585 ± 0.0307	0.515 ± 0.0276	0.663 ± 0.0349
$K_{seg} = C A^{\varphi} L_{seg}^{\xi}$					
C	4.980 ± 0.1683	6.072 ± 0.3184	5.617 ± 0.2916	5.227 ± 0.3136	4.855 ± 0.3650
φ	0.188 ± 0.0109	0.176 ± 0.0185	0.196 ± 0.0182	0.145 ± 0.0171	0.159 ± 0.0223

N number of analysed segments; AL_{seg} , AK_{seg} average segment length and average segment velocity gain computed over participants, frame sizes and trials. L_{seg} , K_{seg} length and velocity gain of individual segments. L_{av} , K_{av} population average of L_{seg} and K_{seg} for each frame size. A area of the frame. Correlations (ρ) and estimates of the parameters (c_0 , c_1 , c_2 , ξ , θ , ψ) of the power-law interpolations obtained by regressing log values of the variables across frame sizes. T_{max}/T_{min} ratio of the axes of the 99% confidence ellipse. C , φ parameters of the predicted relation between A , L_{seg} and K_{seg} . Also indicated the 99% confidence intervals of coefficients and exponents (see Appendix 2)

area, and so did the average segment length (the number of segments was almost constant, irrespective of the frame size, see above), segment lengths were normalized to the population average (L_{av}) for each frame size. Moreover, in order to neutralize the effect of the area on the velocity gain factor, each segment average K_{seg} was normalized to the average gain for the entire trace (K_{av}). Computing the segment gain as $V/R^{*1/3}$, we took into account the fact that the parameter α depends on the average velocity by estimating α for each group and each frame size through the power function approximation derived above.

Figure 11 shows (log–log scale) the relation between the normalized quantities K_{seg}/K_{av} and L_{seg}/L_{av} . The power-function approximation to this relation (Table 3) showed that in all groups the velocity gain factor over a segment was positively correlated with its length (for instance, in young controls $(K_{seg}/K_{av}) = (L_{seg}/L_{av})^{0.162}$). However, a significant difference among groups emerged from the last step of the analysis. Both K_{av} and L_{av} depended on the frame area A . The usual regression analysis in log–log scale showed that a power function provides a good approximation to both relationships (Table 3; for instance, in young controls we found $K_{av} = 5.160 A^{0.258}$; $L_{av} = 1.245 A^{0.428}$). By inserting these approximations in the power-function description of the relation between

K_{seg}/K_{av} and L_{seg}/L_{av} , we finally obtained an expression describing explicitly the way segment gain depended jointly on frame area A and segment length L_{seg} (see Appendix 2). For all groups this procedure for decomposing segment gains yielded

$$\text{Group YC} \quad K_{seg} = 4.980A^{0.188}L_{seg}^{0.162}$$

$$\text{Group AMC1} \quad K_{seg} = 6.072A^{0.176}L_{seg}^{0.169}$$

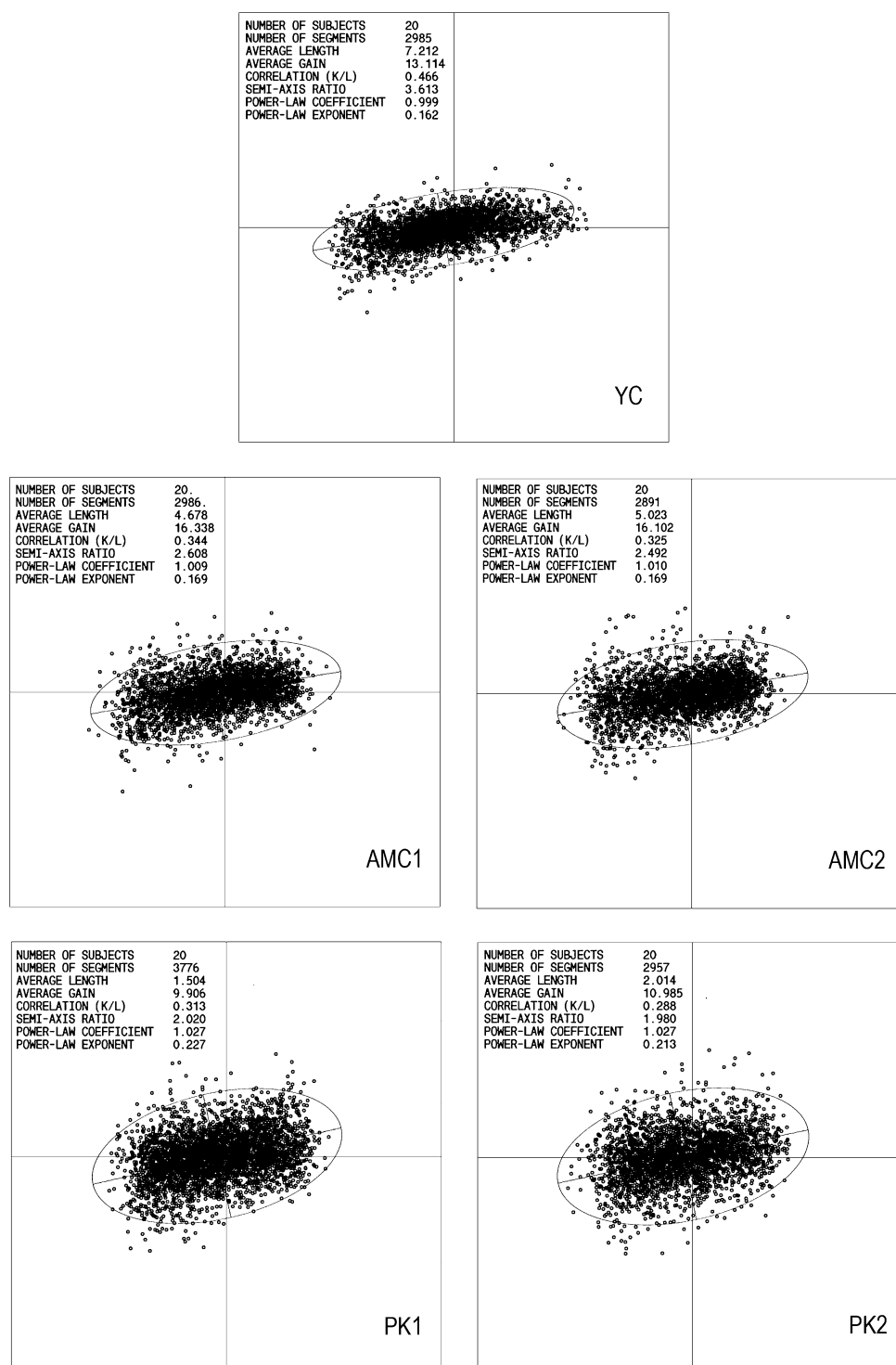
$$\text{Group AMC2} \quad K_{seg} = 5.616A^{0.196}L_{seg}^{0.169}$$

$$\text{Group PRK1} : \quad K_{seg} = 5.227A^{0.145}L_{seg}^{0.227}$$

$$\text{Group PRK2} : \quad K_{seg} = 4.855A^{0.159}L_{seg}^{0.213}$$

Because of the size of the samples involved in the correlation analysis, most 99% confidence intervals for the parameters in the expressions above were small and non-overlapping. In particular, for both exponents, all pair-wise differences between groups were significant ($P < 0.01$). The results, which qualify the analysis of Fig. 9, suggest that the velocity gain factor is set by the joint action of two processes. On the one side, there is a global influence of the frame size captured by the dependence of K_{seg} on the frame area. On the other side, an additional modulation derives from a local form of isochrony that takes into account the length of the segment being executed. This second process

Fig. 11 Local isochrony. Scatter diagrams of the logarithm of the normalized segment length (abscissas) and the logarithm of the normalized velocity gain (ordinates) for each segment of the trajectory. The 99% confidence ellipses are also shown. Results for all groups. The diagrams for the two sessions in groups AMC and PK are shown separately. Full details in the text



is akin to that at work in the case of both single-stroke, and simple closed trajectories. The effect of segment length (gauged by the exponent of the power law; average over groups: 0.188) was weaker than that observed in the latter case (Viviani and Cenzato 1985). However, the main result of our analysis of local isochrony is that the balance

between the weight of local and global isochrony in patients was significantly different from the balance observed in both young and age-matched controls. Specifically, in the former group velocity gain depends more on segment length than frame area. The converse is true for the other two groups.

Discussion

The scribbling traces of the patients revealed the classical cardinal symptoms of PD also observed in other motor tasks. The new finding emerging from our in-depth analysis of velocity control was the dissociation between shape-dependent and shape-independent aspects of the movement.

On the one side, the performance of PD patients exhibited two regularities that are typical of normal hand gestures. First, although the range of values of trace curvature across frame sizes was uniformly higher in patients than in controls (Fig. 5), the traces of the patients were as self-similar across scales as those of the other two groups (Figs. 3, 4). Second, although the traces of the patients were less fluent and regular than those of the control groups (Fig. 2), the Two-thirds Power Law predicted the covariation between instantaneous velocity and curvature with comparable accuracy in both cases (Fig. 7). There are reasons to believe this covariation reflects intrinsic limitations in the neuronal dynamics within the motor cortex (Pellizzer 1997). Thus, the fact that the same instantaneous modulation of velocity was present also in patients suggests that PD has little impact on this dynamics.

On the other side, there was a definite pattern of differences between patients and controls insofar as the timing of the movement is concerned. As concerns velocity, a distinction is in order between absolute and context-dependent control. Confronted with the task of tracing once a given trajectory, everyone chooses idiosyncratically the average velocity of the gesture, largely on the basis of his/her motor habits. All control participants took advantage of this freedom by selecting different baseline tempos for scribbling. Moreover, bradykinesia clearly affected the baseline tempo of patients. Across frame sizes, their average velocity was about 70% lower than that of the other two groups (Fig. 8). However, individual variability was not significantly different among groups.

Context-dependent control of velocity emerges instead when the same gesture is executed successively at different sizes. In this case there is a well-documented spontaneous tendency to modulate the average velocity as a function of movement extent (isochrony). At a global level of analysis (Fig. 8), context-dependent control appeared to be similar in all groups. Although the coefficient of the power-law fit to the frame size/average velocity relation was lowest in patients because of bradykinesia ($YC = 3.03$; $AMC = 3.30$; $PK = 2.28$), the exponents that estimate the strength of the dependence were almost identical ($YC = 0.389$; $AMC = 0.384$; $PK = 0.380$).

By contrast, significant differences among groups were detected by factoring velocity as prescribed by the Two-thirds Power Law (see Fig. 9 and “Results”). The average

velocity gain K across frame sizes was similar among groups ($YC = 13.22$; $AMC = 15.66$; $PK = 12.57$). However, the balance between factors was significantly altered in patients with respect to controls. In young controls the coefficient of the power-law fit to the relation between K and frame size was lowest (5.69) and the exponent was highest (0.236), whereas the opposite was true in patients (average over PK1 and PK2: 6.38 and 0.192, respectively). In other words patients did not scale the gain factor K with the frame size as effectively as controls.

Compared with previous studies of isochrony in PD patients (Van Gemmert et al. 1999, 2003), our results show some significant quantitative differences. The extent to which patients under-scaled movement size (about 70% over a ninefold increase of the frame radius) was similar to that reported by Van Gemmert et al. (2003, Figure 2), but over a much smaller (fivefold) increase of stroke size. More importantly, the range of isochronic behaviour was found to be similar in patients and controls only for very short strokes (up to 2 cm; Van Gemmert 1999). Beyond that limit, automatic velocity compensation in patients was much less effective in patients than in controls. As judged from the peak acceleration (velocity data not available), the relative increase from 1 to 5 cm strokes (Van Gemmert et al. 2003, Figure 3) was strongly different in the two groups (60 and 130%, respectively). Instead, in our study the analogous figures for the average velocity over a wider size range were much higher (437 and 452%), and fairly comparable.

The distinction introduced between *local* and *global* isochrony exposed a further peculiarity of the PK group. The distinction was motivated by the observation that the velocity gain factor K is not actually constant throughout the movement. Whereas the mean gain factor correlates with a *global* size parameter—the area of the frame A —the modulation of K around its mean was correlated with a *local* size parameter, namely the linear extent L_{seg} of the segment of trajectory that is about to be traced. We argue that this modulation reflects the same compensatory mechanism at work when one traces simpler learned trajectories, such as letters in handwriting. Thus, we derived a power-law expression of the gain factor K over a segment in which the contribution of global and local factors is estimated separately (see “Results” *A local form of isochrony*). A contrast between young controls and patients emerged from this analysis, which was particularly clear for the session where patients were in an OFF state (PK1). The exponent of the power law gauging the strength of the global isochrony factor was significantly higher for group YC than for session PK1 (0.188 vs. 0.145). The converse was true for the exponent gauging the strength of the local isochrony factor L_{seg} (0.162 vs. 0.213). Actually, the ratio between factor strengths for YC was almost twice as large

as the ratio for PK1 ($0.188/0.162 = 1.16$ vs. $0.145/0.213 = 0.64$). Two hypotheses may be evoked for the balance shift in patients between local and global compensation. One possible factor is an altered perception of the frame size. In other words patients might underestimate the extent of the space available for the movement. We are not aware of any independent evidence of such a deficit in the perception of space. However, recent work (Desmurget et al. 2000, 2004a) failed to detect a similar deficit in perceived distance. Alternatively, the reduced ability by patients to scale velocity as a function of frame size might reflect the strategic choice to focus attention on the microstructure of the movement to compensate for the reduced accuracy of their gestures.

In the next two sections the dissociation between shape-dependent aspects of the performance that are not affected by PD and shape-independent aspects that are affected is taken up again within the context of current theorizing on the parameters of the movement that are more directly controlled by the motor system.

The control of amplitude and direction of voluntary movements

In considering the role of cortical and sub-cortical centres in motor control, we endorse the so-called *vectorial hypothesis* that direction and amplitude are the relevant parameters for controlling goal-directed movements. The hypothesis is strongly supported by recent behavioural studies (Bock and Eckmiller 1986; Favilla et al. 1990; Bock and Arnold 1992; Gordon et al. 1994a; Pine et al. 1996; Vindras and Viviani 1998), and is consistent with neurophysiological studies showing that movement *direction* is predicted reliably by the tuning curves of the neurons in the motor cortex (Georgopoulos et al. 1983a; Moran and Schwartz 1999). Note that, although the vectorial hypothesis has been formulated mostly on the basis of the results obtained in pointing tasks, it can equally well account for the control of direction in drawing tasks (Schwartz 1992, 1994; Schwartz and Moran 2000).

Demonstrating that movement *amplitude* can also be predicted by cortical activity has proved elusive. The neuronal activity in the primary motor area during a pointing task was found to depend on target distance (Schwartz and Georgopoulos 1987), but the correlation was weak. A later study failed to detect any systematic relation between movement amplitude and the activity in both motor cortex and parietal area 5 (Georgopoulos 1990). This negative result was confirmed by the analysis of wrist flexions and extensions in a reaction time task (Riehle and Requin 1989). Evidence of amplitude coding has been reported in monkeys performing visually guided arm movements (Fu et al. 1993), but reliable correlations were

found only in too few cells to be able to predict movement distance with the same accuracy as movement direction. Other studies also failed to provide convincing evidence that movement amplitude is fully controlled by motor cortex, either because the distance-related modulation of the firing rate occurred after the peak of velocity (Fu et al. 1995), or because of methodological problems (Kurata 1993).

In BG the pattern of correlations between neural activity and movement direction is similar to that found in the motor cortex (Georgopoulos et al. 1983b; Turner and Anderson 1997). In contrast, the search for correlations with movement amplitude has again proved inconclusive. Activity related to movement amplitude and velocity has been observed in a large proportion of subthalamic and pallidal neurons either after movement onset or, less frequently, between the onset of EMG activity and movement onset. Moreover, inhibition or destruction of neurons in the GP reduces the velocity of arm movements (Mink and Thach 1991c; Inase et al. 1996; Desmurget and Turner 2008). However, other studies concluded that BG activity does not relate to amplitude (Brochie et al. 1991; Mink and Thach 1991a, b). Even in the case of simple flexions and extensions of the wrist, the activity of pallidal neurons is strongly dependent on the nature of the task (Mink and Thach 1991a), but quite independent of the kinematic parameters (Mink and Thach 1991b).

A multiplicative model

To reconcile the notion that hand movements are planned in terms of direction and amplitude with the absence of clear neuronal correlates of amplitude coding, Vindras and Viviani (2002) suggested that amplitude is specified jointly by the motor cortex and the BG. Therefore, the activity in either one of these two structures alone is insufficient to account for the final output to the muscles. Here we argue that a further qualification of our hypothesis provides an appropriate framework for discussing the results of the present study.

First, we assume that the motor cortex specifies the spatio-temporal template of the movement, i.e. a blueprint in which both duration and size are free parameters. In the case of scribbling, where no global plan is available before movement begins, partial blueprints are generated sequentially, drawing presumably from a set of basic templates. Second, we also assume that the BG modulate this structural information by taking into account both the entire gesture of which the movement is a component and the context within which the gesture is executed. The assumption is in keeping with the role of context recognition that has been proposed for the BG (Houk and Wise 1995), with the task-related activity observed in pallidal

neurons (Mink and Thach 1991a), and with the suggestion by Turner et al. (1998) that the BG controls the scaling of arm movements.

The modulatory effect of the BG may take place through their projections back to the cortex. Moreover, the BG may also scale the movement by biasing the spinal reflex circuits via its descending output to the brainstem (Johnson et al. 1991; Baldissera et al. 1994; Munro-Davies et al. 1999; Braak et al. 2000; Delwaide et al. 2000; Meunier et al. 2000). The latter hypothesis has a neuro-anatomical basis inasmuch as most axons projecting from the GPi to the thalamus send collaterals to a brainstem area projecting to the reticulo-spinal motor system (Mink 1996).

The simplest implementation of the general framework outlined is suggested by a recent review (Ashe 1997) showing that the slope of the relation between the level of cortical activity and the level of static force is inversely correlated with the range of forces required by the task. Thus, a fixed level of activity may correspond to a whole range of force output, as if a multiplicative factor scaled the effects of the cortical commands on muscles according to specific task requirements. Along the same line of reasoning we assume that the BG modulation action takes the form of a *multiplicative* gain factor acting on both size and velocity.

As for size, form-preserving scaling can be remarkably effective in normal individuals, even when the scale change entails the activation of different body segments (Wiesendanger 1998). Although the control of absolute size is problematic in patients, the self-similarity of the trajectories across frame sizes suggests that the mechanisms underlying context-dependent size scaling are relatively spared by PD. Note that, within the logic of our framework, size scaling must precede velocity scaling because the phenomenon of isochrony implies that average velocity depends on the estimated spatial extent of the movement.

As for the control of velocity, we surmise that the scaling factor is captured by the coefficient K in the Two-thirds Power Law. If so, our analysis of isochrony suggests two conclusions. First, the general reduction of the base tempo in patients is compounded with a reduced ability to scale average velocity as a function of the spatial context (frame size). Second, at the level of single segments this global scaling deficit is partly compensated by a significant increase of the local modulation of the gain factor. In turn, such compensation may reflect an altered balance between the respective roles of cortical and sub-cortical centres.

The fact that the relational aspects of the movement—namely the self-similarity of the trajectories across frame sizes (Figs. 3, 4) and the dependence of the instantaneous velocity on the curvature (Figs. 6, 7)—were rather similar across groups suggests that PD symptoms do not signal a

cortical planning deficit. Instead, the clear evidence of micrographia, and the irregularities often present in the traces of the patients may result from the insufficient size scaling of the movement blueprint compounded with a defective velocity scaling. One cannot rule out that the irregularities are also due to a defective control of the spinal mechanisms that permit fast alternations of muscle contractions and relaxations.

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Appendix 1

We define the parameters selected for the analysis of the movements and detail the data processing for their estimation.

Scaling and smoothing

Traces were aligned by computing the centre of gravity of the samples and shifting all samples so that the new centre was at the origin of the coordinate system. Most of the parameters involve the computation of time derivatives. Because we needed explicit expressions for derivatives (see later), we adopted an interpolation method based on harmonic analysis. The coordinates $x = x(t)$ and $y = y(t)$ of the movement were decomposed in Fourier series:

$$x(t) = \frac{1}{2}a_{x0} + \sum_{k=1}^{\infty} (a_{xk} \cos(k\omega_0 t) + b_{xk} \sin(k\omega_0 t))$$

$$y(t) = \frac{1}{2}a_{y0} + \sum_{k=1}^{\infty} (a_{yk} \cos(k\omega_0 t) + b_{yk} \sin(k\omega_0 t))$$

Preliminary tests showed that retaining the first 50 terms of the series yields an excellent approximation to the traces and is also effective for eliminating uncorrelated noise from the data. All further processing was applied to the truncated series. First and second time derivatives were computed analytically.

Computing the characteristic parameters of the trace

The following parameters were computed from each trace:

(a) *Tangential velocity* [$V(t)$]:

$$V(t) = \sqrt{\left(\frac{dx}{dt}\right)^2 + \left(\frac{dy}{dt}\right)^2}$$

The average velocity of the trace V_0 is related to the total trace length (L) by the equation $V_0 = L/T$.

- (b)
- Total length (L)*
- :

$$L = \int_0^T V(t) dt = \int_0^T \sqrt{\left(\frac{dx}{dt}\right)^2 + \left(\frac{dy}{dt}\right)^2} dt$$

This parameter was computed by integrating numerically the tangential velocity.

- (c)
- Number of inflections [N_I]*
- . An inflection in a 2D trajectory is a point where the curvature of the trajectory changes sign. Inflections were located by identifying the sample index
- k
- such that the quantity

$$\frac{dx(t_k)}{dt} \frac{d^2y(t_k)}{dt^2} - \frac{d^2x(t_k)}{dt^2} \frac{dy(t_k)}{dt}$$

changes sign between k and $k + 1$.

- (d)
- Length of trajectory segments (L_{seg})*
- . Segments were defined as portions of the trajectory bounded by two successive points of inflection. Thus, in a trace there were
- $N_s = N_I - 1$
- segments. When trajectory was almost straight, several inflections occurred in close succession.

- (e)
- Time average of the one-third power of the radius [R_m^{1/3}]*
- . The function
- $R(t)$
- that describes how the radius of curvature of the trajectory changes in time is

$$R(t) = \frac{V(t)^3}{\left| \frac{dx}{dt} \frac{d^2y}{dt^2} - \frac{dy}{dt} \frac{d^2x}{dt^2} \right|}$$

The radius of curvature increases in the proximity of a point of inflection where it becomes infinite. The most recent formulation of Two-thirds Power Law circumvents this difficulty by expressing the velocity as a power function of the normalized radius $R^*(t)$ defined as

$$R^*(t) = \frac{R(t)}{1 + \alpha R(t)}$$

As the movement approaches an inflection, R^* stays finite and tends to the limit $1/\alpha$. The required time average $\langle R^{*1/3} \rangle$ was computed by numerical evaluation of the integral

$$\langle R^{*1/3} \rangle = \frac{1}{T} \int_0^T R^*(t)^{1/3} dt$$

- (f)
- Average curvature [C_m]*
- . Curvature is the inverse of the radius of curvature. It would be inappropriate to compute the average curvature using directly the Fourier approximation of the trajectory. Because the instantaneous velocity decreases with curvature as prescribed by the Two-thirds Power Law, and because

the sampling rate is constant, the sample density around the high-curvature portions of the trajectory is much higher than the density within low-curvature portions. Thus, numerical integration of the inverse of the radius $R(t)$ would severely overestimate the average geometric curvature. Instead, we adopted the following resampling strategy. Let us consider a doubly differentiable time function $\varphi = \varphi(t)$ such that $\varphi(0) = 0$ and $\varphi(T) = T$, where T is the total duration of the movement. The parametric equations $[x = x(\varphi(t)), y = y(\varphi(t))]$ describe the same trajectory C as the original ones $[x = x(t), y = y(t)]$. However, the kinematics of the movement depends on the function φ , and the correspondent transformed velocity is in general different from the velocity of the actual movement:

$$V_\varphi(t) = \frac{d\varphi}{dt} \left[\left(\frac{dx}{d\varphi}\right)^2 + \left(\frac{dy}{d\varphi}\right)^2 \right]^{1/2}$$

The expression above can be rewritten as a separable nonlinear differential equation:

$$d\varphi = V_\varphi(t) \left[\left(\frac{dx}{d\varphi}\right)^2 + \left(\frac{dy}{d\varphi}\right)^2 \right]^{1/2} dt$$

Under mild continuity conditions, for any choice of the transformed velocity function $V_\varphi(t)$, solving the equation above yields the unique function φ that is compatible with this choice. We imposed the condition that the tangential velocity is constant and equal to the average velocity of the actual movement ($V_\varphi(t) = L/V_m$) and computed the solution $\varphi(t)$ with a fourth-order Runge–Kutta algorithm with the boundary conditions $\varphi(0) = 0$ and $\varphi(T) = T$. By inserting the solution $\varphi(t)$ back into the parametric equations, the original movement was resampled so that successive data points were spaced by a constant fraction of the total length rather than by a constant time interval (the total number of samples was kept equal to 2,000 as in the original trace). Finally, the average geometric curvature was calculated as the mean over all samples of the inverse of the radius. Note that this strategy was possible only because the Fourier series affords an analytical approximation to the traces.

- (g)
- Average gain factor [K_m]*
- . According to the
- Two-Thirds Power Law*
- , the multiplicative parameter
- K
- (gain factor) is approximately constant over successive units of motor action. We computed the average gain factor for the entire trace by numerical estimation of the integral

$$K_m = \frac{1}{T} \int_0^T \left| \frac{dx(t)}{dt} \frac{d^2y(t)}{dt^2} - \frac{dy(t)}{dt} \frac{d^2x(t)}{dt^2} \right|^{\frac{1}{3}} dt$$

In the same manner, we computed the average gain factors K_{seg} for each segment within the complete trace.

Appendix 2

We specify the method for testing the statistical significance of parameters in the equation relating frame area A and segment length L_{seg} to segment gain K_{seg} . The three equations to be combined are (relevant stochastic variables in boldface)

$$K_{\text{seg}}/K_{\text{av}} = \mathbf{c}_0 (L_{\text{seg}}/L_{\text{av}})^{\xi}$$

$$K_{\text{av}} = \mathbf{c}_1 A^{\theta}$$

$$L_{\text{av}} = \mathbf{c}_2 A^{\psi}$$

As expected, \mathbf{c}_0 was almost indistinguishable from 1. Therefore

$$K_{\text{seg}} = \mathbf{c}_1 \mathbf{c}_2^{-\xi} A^{(\psi - \theta \xi)} L_{\text{seg}} = \mathbf{C} A^{\varphi} L_{\text{seg}}$$

where $\mathbf{C} = \mathbf{c}_1 \mathbf{c}_2^{-\xi}$ and $\varphi = \psi - \theta \xi$. The variances σ_c^2 of the exponents ξ , θ and ψ are estimated directly from the regression equations

$$\begin{aligned} \log(K_{\text{av}}) &= \log(\mathbf{c}_1) + \theta \log(A); \\ \log(L_{\text{av}}) &= \log(\mathbf{c}_2) + \psi \log(A) \end{aligned}$$

through the formula (Kendall and Stuart 1968, p. 395):

$$\sigma_e^2 = \left(\sigma_y^2 / \sigma_x^2 \right) (1 - \rho^2) / (N - 2)$$

where ρ is Fisher's correlation coefficient and N is the sample size. The averages of the regression coefficients $\mathbf{a}_1 = \log(\mathbf{c}_1)$ and $\mathbf{a}_2 = \log(\mathbf{c}_2)$ are $\mu_{a1} = a_1$ and $\mu_{a2} = a_2$, respectively. Their variances σ_a^2 are estimated by

$$\sigma_a^2 = \sigma_e^2 (\sigma_x^2 [N - 1] / N + \mu_x^2)$$

If two stochastic variables are related by a monotonic function $\mathbf{y} = \mathbf{g}(\mathbf{x})$ the pdf of \mathbf{y} is given by $f_y(\mathbf{y}) = f_x(\mathbf{g}^{-1}(\mathbf{y})) / d\mathbf{g}(\mathbf{g}^{-1}(\mathbf{y})) / d\mathbf{x}$ (Papoulis 1965, p. 126). We assume that both \mathbf{a}_1 and \mathbf{a}_2 have a Gaussian pdf. Thus, because $\mathbf{c}_1 = \exp(\mathbf{a}_1)$ and $\mathbf{c}_2 = \exp(\mathbf{a}_2)$ the pdf of \mathbf{c}_1 and \mathbf{c}_2 have the common expression

$$\frac{\exp\left(-\frac{(\log(c) - \mu_c)^2}{2\sigma_c^2}\right)}{c \sqrt{2\pi\sigma_c^2}}$$

By computing first and second moments of this distribution, average and variance of \mathbf{c}_1 and \mathbf{c}_2 can be expressed in terms of known quantities

$$\mu_c = \exp(\mu_a + \sigma_a^2/2)$$

$$\sigma_c^2 = \exp(\mu_a + 2\sigma_a^2) - \exp(\mu_a + \sigma_a^2)$$

Finally, the pdf of $\mathbf{C} = \mathbf{c}_1 \mathbf{c}_2^{-\xi}$ and $\varphi = \theta - \psi \xi$ cannot be computed in closed form. However, the pdf of all stochastic variables appearing in the expressions of \mathbf{C} and φ are known. Thus, the variances σ_C^2 and σ_φ^2 and the corresponding 99% confidence intervals were finally estimated with a Montecarlo procedure ($n = 30,000$).

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